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UTILITY
PATENT APPLICATION
TRANSMITTAL

Attorney Docket No. 7969-087 Total Pages 146

First Named Inventor or Application Identifier

W. James Jackson

	P) Hy for new nonprovisional applications under 37 CFR 1.53(b))	Express	Mail Label No. EL 501 634 175 US
APPLICATION ELEMENTS See MPEP chapter 600 concerning utility patent application contents.			Assistant Commissioner for Palassistant Commissioner for Palassistant ADDRESS TO: Box Patent Application Washington, DC 20231
1. 🗵	Fee Transmittal Form Submit an original, and a duplicate for fee processing)		6. Microfiche Computer Program (Appendix)
2. 🛚	Specification [Total Page (preferred arrangement set forth below)	s <u>89</u>]	7. ⋈ Nucleotide and/or Amino Acid Sequence Submission (if applicable, all necessary)
	-Table of Contents -Descriptive title of the Invention		a. 図 Computer Readable Copy
	-Cross Reference to Related Applications		b. 図 Paper Copy (identical to computer copy)
	-Statement Regarding Fed sponsored R&D		
	-Reference to Microfiche Appendix		c. Statement verifying identity of above copies
	-Background of the Invention	Γ	A DELICATION DADTO
	-Brief Summary of the Invention		ACCOMPANYING APPLICATION PARTS
	-Brief Description of the Drawings (if filed) -Detailed Description of the Invention (including drawings, if filed)		8. Assignment Papers (cover sheet & document(s))
	-Claim(s)		9. ☐ 37 CFR 3.73(b) Statement ☐ Power of Attorney (when there is an assignee)
7d	-Abstract of the Disclosure - Drawing(s) (35 USC 113)	ote 14.1	10. □ English Translation Document (if applicable)
3: ≥ 4. ⊠	ITatal Char		11. □ Information Disclosure □ Copies of IDS Statement (IDS)/PTO-1449 Citations
	a. Newly executed (original or copy)		12. Preliminary Amendment
4.42	Newly executed (original of copy) Copy from a prior application (37 CFR 1.63(d)) (for continuation/divisional with Box 17 completed)		13. Return Receipt Postcard (MPEP 503) (Should be specifically itemized)
ļ-L	[Note Box 5 below]		14. ≅ Small Entity □ Statement filed in prior application, Statement(s) Status still proper and desired
	i. DELETION OF INVENTORS(S)	the prior	15. □ Certified Copy of Priority Document(s)
	Signed statement attached deleting inventor(s) named in tapplication, see 37 CFR 1.63(d)(2) and 1.33 (b).	rie prior	(if foreign priority is claimed)
	Incorporation By Reference (useable if Box 4b is checked)	of the oath	16. DOther:
	The entire disclosure of the prior application, from which a copy or declaration is supplied under Box 4b, is considered as being p	art of the	
1000	disclosure of the accompanying application and is hereby incorporate	orated by	
	reference therein.	-1.46	ita information:
17.	If a CONTINUING APPLICATION, check appropriate box and support of the continuation □ Divisional □ Continuation-in-part (CIP)	of prior a	application No. Tiled.
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ATTORNEY DOCKET NO. 7969-087

Date: October 2, 2000

Assistant Commissioner for Patents Box PATENT APPLICATION Washington, D.C. 20231

Sir:

The following utility patent application is enclosed for filing:

Executed on: October 2, 2000 Applicant(s): W. James Jackson

CHLAMYDIA PROTEIN, GENE SEQUENCE AND USES THEREOF Title of Invention:

PATENT APPLICATION FEE VALUE

ТҮРЕ	NO. FILED	LESS	EXTRA	EXTRA RATE	F	EE
Total Claims	104	-20	84	\$18.00 each	\$	1,512.00
Independent	8	-3	5	\$80.00 each	\$	400.00
Application of the state of the			Minimum Fee		\$	690.00
	Multiple Dependency Fee If Applicable (\$270.00)					270.00
The state of the s			Total		\$	2,872.00
Harry Brazil	50% Reduction for Independent Inventor, Nonprofit Organization or Small Business Concern (a verified statement as to the applicant's status is attached) - \$ 1,453.00					
400 to 100 to 10				Total Filing Fee	\$	1,419.00

Priority of application no. filed on in is claimed under 35 U.S.C. § 119.

The certified copy of the priority application has been filed in application no. filed

Amend the specification by inserting before the first line the following sentence: This is a continuation-in-part of application no. filed.

Please charge the required fee to Pennie & Edmonds LLP Deposit Account No. 16-1150. A copy of this sheet is

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Respectfully submitted,

Geraldine F. Baldwin

PENNIE & EDMONDS LLP

Enclosure

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Express Mail No.: <u>EL 501 634 175 US</u>

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Application of: W. James Jackson

Application No.: To Be Assigned

Group Art Unit:

Filed: Concurrently Herewith

Examiner:

For:

CHLAMYDIA PROTEIN, GENE

Attorney Docket No.: 7969-087

SEQUENCE AND USES THEREOF

REQUEST TO ESTABLISH SMALL ENTITY STATUS AND TRANSMITTAL OF VERIFIED STATEMENT CLAIMING SMALL ENTITY STATUS UNDER 37 C.F.R. § 1.9(f) AND 1.27(c)(d)

Assistant Commissioner for Patents Washington, D.C. 20231

Sir:

Attorneys for Applicant request that Small Entity Status be established for the above-identified application and submit herewith an original executed Verified Statement Claiming Small Entity Status - Small Business Concern from Antex Biologics, Inc., assignee of the above-identified application by virtue of assignment from the inventor W. James Jackson filed herewith. It is requested that this document be made of record in the file of the above-identified application.

Respectfully submitted,

Date October 2, 2000

Geraldine F. Baldwin (Reg. No.)

PENNIE & EDMONDS LLP 1155 Avenue of the Americas New York, New York 10036-2711

(212) 790-9090

Enclosure

Dy Margaret B. Bridanlan Reg No. 40,922

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re:

□ Patent of:

■ Application No.: To Be Assigned

Group Art Unit:

☐ Patent No.:

Examiner:

□ Issued:

For:

CHLAMYDIA PROTEIN, GENE

SEQUENCE AND USES THEREOF

Attorney Docket No.: 7969-087

VERIFIED STATEMENT (DECLARATION) CLAIMING SMALL ENTITY STATUS [37 CFR 1.9(f) and 1.27(c)] - Small Business Concern

Assistant Commissioner for Patents Washington, D.C. 20231

Sir:

I hereby declare that I am

□ the owner of the small business concern identified below:

an official of the small business concern empowered to act in behalf of the concern identified below:

Name of organization Antex Biologics, Inc.

Address of organization 300 Professional Drive

Gaithersburg, Maryland 20879

I hereby declare that the above identified small business concern qualifies as a small business concern as defined in 37 CFR 1.9(d), for purposes of paying reduced fees under section 41(a) and (b) of Title 35, United States Code, in that the number of employees of the concern, including those of its affiliates, does not exceed 500 persons. For purposes of this statement, (1) the number of employees of the business concern is the average over the previous fiscal year of the concern of the person employed on a full-time, part-time or temporary basis during each of the pay periods of the fiscal year, and (2) concerns are affiliates of each other when either, directly or indirectly, one concern controls or has the power to control the other, or a third party or parties controls or has the power to control both.

I hereby declare that rights under contract or law have been conveyed to and remain with the small business concern and/or there is an obligation under contract or law by the inventor(s) to convey rights to the small business concern with regard to the invention entitled CHLAMYDIA PROTEIN, GENE SEQUENCE AND USES THEREOF by inventor(s) W. James Jackson described in

the specification filed herewith

application no. filed

patent no. issued

NY2 - 1126269.1

If the rights held by the above identified small business concern are not exclusive, each individual, concern or organization having rights to the invention is listed below and no rights to the invention are held by any person, other than the inventor, who could not qualify as an independent inventor under 37 CFR 1.9(c) if that person made the invention, or by any concern which would not qualify as a small business concern under 37 CFR 1.9(d), or a nonprofit organization under 37 CFR 1.9(e).

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□ INDIVIDUAL	□ SMALL E	BUSINESS CONCERN	□ NONPR	OFIT ORGANIZATION	
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(37 CFR 1.27)

CEO

300 Professional Drive Gaithersburg, Maryland 20879

Title of person other than owner Address of person signing

Signature

and Cheirman

Date

NY2 - 1126269.1

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CHLAMYDIA PROTEIN, GENE SEQUENCE AND USES THEREOF



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CHLAMYDIA PROTEIN, GENE SEQUENCE AND USES THEREOF

1. FIELD OF THE INVENTION

The present invention generally relates to PMP polypeptides of *Chlamydia*, amino acid and nucleotide sequences thereof, antibodies specific for such *Chlamydia* PMP polypeptides, prophylactic and therapeutic compositions, including vaccines, and to methods of preventing, treating or ameliorating disorders in mammals and birds related to *Chlamydia* infections and for inducing immune responses in animals to *Chlamydia*.

10 2. BACKGROUND OF THE INVENTION

Chlamydiae are obligate intracellular bacteria that infect animals, including mammals and birds, particularly at the epithelial lining of the lung, conjunctivae or genital tract. The most common species of Chlamydia include Chlamydia trachomatis, Chlamydia psittaci, Chlamydia pecorum and Chlamydia pneumoniae. Recently, the newly designated species of Chlamydia, C. pneumoniae (formerly C. trachomatis TWAR), has been implicated as a major cause of epidemic human pneumonitis and perhaps may play a role in atherosclerosis.

There are currently 18 recognized *C. trachomatis* serovars, causing trachoma and a broad spectrum of sexually transmitted diseases, with the A, B and C serovars being most frequently associated with trachoma, while the D-K serovars are the most common cause of genital infections.

Chlamydia are prevalent human pathogens causing disorders such as sexually transmitted diseases, respiratory diseases, including pneumonia, neonatal conjunctivitis, and blindness. Reactive inflammatory arthritis is a common sequel to sexually acquired non-gonococcal genital tract infection. Approximately 50% of reactive inflammatory arthritis cases are associated with Chlamydia trachomatis infection of the genital tract.

C. trachomatis is the major cause of sexually transmitted disease in many industrialized countries, including the United States. While the exact incidence of C.
 30 trachomatis infection in the United States is not known, current epidemiological studies indicate that more than 4 million chlamydial infections occur each year, compared to an estimated 2 million gonococcal infections. While all racial, ethnic and socioeconomic groups are affected, the greatest number of chlamydial infections occurs among young, 12 to 20 year-old, sexually active individuals. Most genitourinary chlamydial infections are
 35 clinically asymptomatic. Prolonged carriage in both men and women is common. As many as 25% of men and 75% of women diagnosed as having chlamydial infections have no overt

signs of infection. As a consequence, these asymptomatic individuals constitute a large reservoir that can sustain transmission of the agent within the community.

Far from being benign, serious disease can develop from these infections including: urethritis, lymphogranuloma venereum (LGV), cervicitis, and epididymitis in males. Ascending infections from the endocervix commonly gives rise to endometritis, pelvic inflammatory disease (PID) and salpingitis which can cause tubal occlusion and lead ultimately to infertility in females.

C. trachomatis infection of neonates results from perinatal exposure to the mother's infected cervix. Nearly 70% of neonates born vaginally to mothers with
10 chlamydial cervicitis become infected during delivery. The mucus membranes of the eye, oropharynx, urogenital tract and rectum are the primary sites of infection. Chlamydial conjunctivitis has become the most common form of ophthalmia neonatorum.
Approximately 20-30% of exposed infants develop inclusion conjunctivitis within 14 days of delivery even after receiving prophylaxis with either silver nitrate or antibiotic ointment.
15 C. trachomatis is also the leading cause of infant pneumonia in the United States. Nearly

10-20% of neonates delivered through an infected cervix will develop chlamydial pneumonia and require some type of medical intervention.

In developing countries, ocular infections of *C. trachomatis* cause trachoma,

a chronic follicular conjunctivitis where repeated scar formation leads to distortion of the eyelids and eventual loss of sight. Trachoma is the world's leading cause of preventable blindness. The World Health Organization estimates that over 500 million people worldwide, including about 150 million children, currently suffer from active trachoma and over 6 million people have been blinded by this disease.

In industrialized countries, the costs associated with treating chlamydial infections are enormous. In the United States, the annual cost of treating these diseases was estimated at \$2.5-3 billion in 1992 and has been projected to exceed \$8 billion by the year 2000.

One potential solution to this health crisis would be an effective chlamydial vaccine. Several lines of evidence suggest that developing an effective vaccine is feasible.

Studies in both humans and primates have shown that short-term protective immunity to *C. trachomatis* can be produced by vaccinating with whole *Chlamydia*. However, protection was characterized as short lived, serovar specific, and due to mucosal antibody production. Additionally, in some vaccinees disease was exacerbated when these individuals became naturally infected with a serovar different from that used for

35 immunization. This adverse reaction was ultimately demonstrated to be due to a delayed-

type hypersensitivity response. Thus, the need exists to develop a subunit-based chlamydial vaccine capable of producing an efficacious but nonsensitizing immune response. Such a subunit vaccine may need to elicit both mucosal neutralizing secretory IgA antibody and/or cellular immune response to be efficacious.

Subunit vaccine development efforts to date have focused almost exclusively on the major outer membrane protein (MOMP). MOMP is an integral membrane protein of approximately 40 kDa in size and comprises up to about 60% of the infectious elementary body (EB) membrane protein (Caldwellet al. 1981. Infect. Immun., 31:1161-1176). MOMP imparts structural integrity to the extracellular EB and is thought to function as a porin-like 10 molecule when the organism is growing intracellularly and is metabolically active. With the exception of four surface exposed variable domains (VDI-VDIV), MOMP is highly conserved among all 18 serovars. MOMP is highly immunogenic and can elicit a local neutralizing anti-Chlamydia antibody. However, problems exists with this approach.

To date, most MOMP-specific neutralizing epitopes that have been mapped 15 are located within the VD regions and thus give rise only to serovar-specific antibody. Attempts to combine serovar-specific epitopes in various vaccine vectors (e.g., poliovirus) to generate broadly cross-reactive neutralizing antibodies have been only marginally successful (Murdin et al. 1993. Infect. Immun., 61:4406-4414; Murdin et al. 1995. Infect. Immun., 63:1116-1121).

20 Two other major outer membrane proteins in C. trachomatis, the 60 kDa and 12 kDa cysteine-rich proteins, as well as the surface-exposed lipopolysaccharide, are highly immunogenic but, unlike MOMP, have not been shown to induce a neutralizing antibody (Cerrone et al., 1991, Infect. Immun., 59:79-90). Therefore, there remains a need for a novel subunit-based chlamydial vaccine.

25 Citation or identification of any reference in this section or any other section of this application shall not be construed as an indication that such reference is available as prior art to the present invention.

3. SUMMARY OF THE INVENTION

30 This invention is directed to PMP (putative membrane proteins) proteins (referred to hereafter and in the claims as PMP polypeptides or PMP proteins) from Chlamydia spp. More particularly, the present invention encompasses the family of PMPE and PMPI polypeptides of Chlamydia trachomatis and other Chlamydia spp., including but not limited to, Chlamydia pneumonia, Chlamydia pecorum, and Chlamydia psittaci, having 35 a molecular weight of 90 to 115 kD, and an amino acid sequence of or homologous to SEQ

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ID NO.:2 (PMPE) or SEQ ID NO.:4 (PMPI), in isolated or recombinantly produced form. SEQ ID NOs.:2 and 4 represent the amino acid sequences of the *Chlamydia trachomatis* L2 serovar PMPE and PMPI proteins, respectively, encoded by the *pmpE* and *pmpI* genes. The nucleotide sequences for the *pmpE* and *pmpI Chlamydia trachomatis* L2 serovar coding regions are SEQ ID NOs.:1 and 3, respectively. The present invention encompasses all *Chlamydia* PMPE and PMPI polypeptides, particularly those of the *Chlamydia trachomatis* L2 serovar, and also including PMPI and PMPE polypeptides from other *Chlamydia trachomatis* serovars and other *Chlamydia* species. Identification of these homologs and analogs can be accomplished by methods well known in the art, for example, but not limited to, nucleic acid hybridization and PCR based techniques. The present invention encompasses a purified PMPE or PMPI polypeptide, polypeptides derived therefrom ("PMP-derived polypeptides", *e.g.*, derivatives, fragments and analogs thereof), preferably, that elicit an immune reaction against whole *Chlamydia* cells and/or are bound by anti-PMPE or anti-PMPI antibodies, and methods for making said polypeptides and PMP-derived polypeptides.

Preferably the PMP protein has the amino acid sequence of SEQ IN NO.:2 or 4 or is substantially homologous to any of SEQ ID NO.:2, 4 or 5-34. Preferred fragments of the protein comprise an amino acid sequence of any of SEQ ID NOs.:5-34.

Preferably, the PMP protein is an outer membrane protein. More preferably, the PMP protein is surface localized. Preferably, the PMP protein has at least one GGAI (Gly Gly Ala Ile) domain. It is intended that PMP proteins from all species of *Chlamydia* are included in this invention; however, preferred species include *Chlamydia trachomatis*, *Chlamydia psittaci*, *Chlamydia pecorum* and *Chlamydia pneumoniae*.

The invention also provides PMP fusion peptides having B and/or T-cell stimulating activity, preferably comprising at least two T or B cell epitopes derived from the same or from different *Chlamydia* PMP proteins which proteins, or portions thereof, are arranged in a contiguous polypeptide in a configuration different from a naturally occurring configuration of the regions of a *Chlamydia* PMP protein.

A preferred polypeptide of the invention is a fusion polypeptide comprising at least two peptides, said at least two peptides consisting of amino acid sequences selected from the group consisting of SEQ ID NOs.:5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17,18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, and 34, with the proviso that the peptides of the polypeptide are arranged in a contiguous polypeptide configuration that is different from the configuration of a naturally occurring PMPE or PMPI polypeptide (*e.g.*,

35 is not a naturally occurring PMPE or PMPI polypeptide).

Other preferred PMP-derived polypeptides of the invention are isolated fusion polypeptides wherein the polypeptide comprises an amino acid sequence of SEQ ID NO.:5, 6, 7, 8, 9, 10 or 11 or an isolated fusion polypeptide wherein the polypeptide comprises an amino acid sequence of SEQ ID NO.:23, 24, 25, 26, 27, 28, or 29, with the proviso that the peptides of the polypeptide are arranged in a configuration that is different from the configuration of a naturally occurring PMPE or PMPI polypeptide. A preferred PMP-derived polypeptide is an isolated fusion polypeptide, wherein the polypeptide comprises an amino acid sequence of SEQ ID NO.:5, 6, 7, 8, 9, 10, 11, 23, 24, 25, 26, 27, 28 or 29, with the proviso that the peptides are arranged in a configuration that is different from the configuration of a naturally occurring PMPE or PMPI polypeptide.

Preferably, the PMP-derived polypeptides of the invention are immunologically cross-reactive with the PMP protein from which they are derived and are capable of eliciting in an animal an immune response to *Chlamydia*. A preferred PMP polypeptide or PMP-derived polypeptide of the invention induces IgM, IgG, IgA, and/or IgE antibodies, delayed hypersensitivity T-cell responses and/or cytotoxic T-cell responses to cells expressing Chlamydial antigens (including but not limited to cells infected with *Chlamydia* and antigen presenting cells such as macrophages, dendritic cells, B cells, or synthetic antigen presenting cells which display Chlamydial antigens), native PMP protein from which the polypeptide is derived, *Chlamydia* cells, or *Chlamydia* elemental bodies (EB). In a more preferred embodiment, the PMP polypeptide or PMP-derived polypeptide is capable of eliciting an immune response against other *Chlamydia* serovars and, more preferably, other *Chlamydia* species along with the *Chlamydia* serovar or species in which

The invention also encompasses antisera and antibodies, including but not limited to, cytotoxic or bactericidal polyclonal and monoclonal antibodies, which bind to and are specific for the PMP polypeptide, PMP-derived polypeptides and/or fragments thereof.

the PMP polypeptide occurs naturally.

Preferably the antibodies bind a PMP protein having the amino acid sequence of SEQ ID NOs.:2 or 4 or an amino acid sequence substantially homologous thereto. Also included are monoclonal antibodies that specifically bind a PMP or PMP-derived polypeptide, including but not limited to monoclonal antibodies that specifically bind a polypeptide consisting of an amino acid sequence of any of SEQ ID NOs.:2, 4 or 5-34. Also included are antigen binding fragments of polyclonal or monoclonal antibodies, e.g., Fv, Fab, Fab' and F(ab')₂ fragments. A further aspect of the invention are chimerized or humanized antibodies in which one or more of the antigen binding regions of the anti-

PMP antibody is introduced into the framework region of a heterologous (e.g., human) antibody. In a preferred aspect, the antibodies are human antibodies.

Another aspect of the invention is directed to T-cells raised against an antigenic or immunogenic composition of the invention or T-cells specific for antigenic or immunogenic polypeptides of the invention or specific for cells expressing Chlamydial antigens (including but not limited to cells infected with *Chlamydia* or antigen presenting cells presenting PMP polypeptides such as dendritic cells, B cells, or synthetic antigen presenting cells), *Chlamydia* cells, or *Chlamydia* elemental bodies (EB).

The invention further provides isolated nucleic acid molecules (DNA or RNA) encoding the PMPE polypeptides, PMPI polypeptides, PMPE-derived polypeptides, PMPI-derived polypeptides, vectors having said sequences, host cells containing said vectors or having the sequences operably linked to a heterologous promoter, recombinant polypeptides produced therefrom, and pharmaceutical compositions comprising the nucleic acid molecules, vectors, and cells.

A preferred aspect of the invention is a nucleotide sequence encoding a PMP protein comprising the amino acid sequence of any of SEQ ID NOs.:2, 4 or 5-34 or an amino acid sequence substantially homologous thereto. Also included is an isolated nucleic acid molecule comprising a DNA sequence of SEQ ID NOs.:1 or 3 or a complementary sequence thereof; a fragment of the DNA molecule having the nucleotide sequence of SEQ ID NOs.:1 or 3, or the complimentary sequence thereof; or a nucleic acid molecule which hybridizes under stringent conditions to any one of the sequences described above. The nucleic acid molecule that hybridizes under stringent conditions preferably has a sequence homology of about 70%, 80%, 90%, 95%, or 99% with any of the sequences identified above, more preferably about 90%.

The invention further encompasses pharmaceutical compositions including prophylactic or therapeutic compositions, which may be immunogenic compositions, including vaccines, comprising one or more of the PMP polypeptides of the invention, optionally in combination with, fused to, or conjugated to one or more other component(s), including a lipid, phospholipid, a carbohydrate, including a lipopolysaccharide, any proteins either novel or known to those skilled in the art, inactivated whole or attenuated organisms, including but not limited to viruses, yeasts, fungi and bacteria. Particularly preferred bacteria include, but are not limited to *Neisseria*, *Chlamydia*, *Moraxella*, *Pseudomonas*, *Streptococcus* or *Haemophilus* bacteria.

In a specific embodiment, the invention encompasses pharmaceutical compositions, including prophylactic or therapeutic compositions, which may be

immunogenic compositions including vaccines, comprising one or more of the PMP polypeptides and/or PMP-derived polypeptides and an attenuated or inactivated Chlamydia cultivar or an attenuated or inactivated Chlamydia cultivar expressing PMP polypeptide in a greater amount when compared to wild-type Chlamydia.

The invention further encompasses pharmaceutical compositions comprising isolated nucleic acid molecules encoding PMP polypeptides and PMP-derived polypeptides of the present invention which can be used in methods to detect Chlamydia infection or to prevent, treat or reduce the severity of a disease or disorder related to infection with Chlamydia. Such compositions include but are not limited to vectors or recombinant host 10 cells comprising such nucleic acid molecules or having a nucleotide sequence of the invention operably linked to a heterologous promoter.

The invention also includes diagnostic reagents, that may include any one or more of the above mentioned aspects, such as native PMP proteins, recombinant PMP proteins, PMP-derived polypeptides, nucleic acid molecules, immunogenic compositions, 15 antigenic compositions, antisera, T-cells, antibodies, vectors comprising the nucleic acids, and transformed cells comprising the vectors.

A further aspect of the present invention provides methods for determining the presence of nucleic acids encoding a PMP protein or a PMP-derived polypeptide in a test sample, and diagnostic kits and reagents therefor, for determining the presence of a 20 nucleic acid encoding a PMP polypeptide or PMP-derived polypeptide.

Also included in this invention are methods of inducing an immune response to Chlamydia spp. and methods of preventing, treating or ameliorating disorders or diseases related to Chlamydia in an animal, including mammals and birds and, preferably, in humans, in need of such treatment comprising administering an effective amount of the 25 pharmaceutical or vaccine composition of the invention. Preferred disorders or diseases include a Chlamydia bacterial infection, trachoma, conjunctivitis, urethritis, lymphogranuloma venereum (LGV), cervicitis, epididymitis, or endometritis, pelvic inflammatory disease (PID), salpingitis, tubal occlusion, infertility, cervical cancer, reactive arthritis, and atherosclerosis.

30 A further aspect of the invention relates to antagonists or agonists which inhibit or enhance the activity or expression of the polypeptides or nucleic acid molecules of the invention. Preferred are bacteriostatic or bacteriocidal agonists or antagonists.

A further aspect of the invention is a method for identifying compounds which interact with and inhibit or activate an activity of the polypeptides or nucleic acid 35 molecules of the invention comprising contacting a composition comprising the polypeptide or the nucleic acid molecule with the compound to be screened under conditions that permit interaction between the compound and the polypeptide or nucleic acid molecule to assess the interaction of a compound and to detect interaction of the compound with the polypeptide or nucleic acid. The interaction of the compound with the polypeptide or nucleic acid molecule is determined by the association of a second component (e.g., an antibody) capable of providing a detectable signal in response to the interaction of the polypeptide or nucleic acid molecule with the compound; and determining the presence or absence of a signal generated from the interaction of the compound with the polypeptide or nucleic acid molecule. Alternatively, the interaction of the compound to inhibit the activity of the polypeptide or the nucleic acid molecule.

3.1 <u>ABBREVIATIONS</u>

	anti-PMP	= PMP polypeptide antibody or antiserum		
15	ATCC	= American Type Culture Collection		
	immuno-reactive	= capable of provoking a cellular or humoral immune		
		response		
	kD or kDa	= kilodaltons		
	OG	= n-octyl-D-glucopyranoside or octyl glucoside		
20	OMP	= outer membrane protein		
	OMPs	= outer membrane proteins		
	PBS	= phosphate buffered saline		
	PAGE	= polyacrylamide gel electrophoresis amino acid residues		
	polypeptide	= a peptide of any length, preferably having eight or more		
25		amino acid residues		
	SDS	= sodium dodecylsulfate		
	SDS-PAGE	= sodium dodecylsulfate polyacrylamide gel electrophoresis		

Nucleotide sequences defined herein are represented by one-letter symbols

30 for the bases as follows:

A (adenine)
C (cytosine)

G (guanine)

T (thymine)

35 U (uracil)

```
M (A or C)
R (A or G)
W (A or T/U)
S (C or G)

5 Y (C or T/U)
K (G or T/U)
V (A or C or G; not T/U)
H (A or C or T/U; not G)
D (A or G or T/U; not C)

10 B (C or G or T/U; not A)
```

N (A or C or G or T/U) or (unknown)

Peptide and polypeptide sequences defined herein are represented by one-

letter symbols for amino acid residues as follows:

- 15 A (alanine)
 - R (arginine)
 - N (asparagine)
 - D (aspartic acid)
 - C (cysteine)
- 20 Q (glutamine)
 - E (glutamic acid)
 - G (glycine)
 - H (histidine)
 - I (isoleucine)
- 25 L (leucine)
 - K (lysine)
 - M (methionine)
 - F (phenylalanine)
 - P (proline)
- 30 S (serine)
 - T (threonine)
 - W (tryptophan)
 - Y (tyrosine)
 - V (valine)
- 35 X (unknown)

The present invention may be more fully understood by reference to the following detailed description of the invention, non-limiting examples of specific embodiments of the invention and the appended figures.

5 4. BRIEF DESCRIPTION OF DRAWINGS

Figure 1. Schematic map of the *C. trachomatis* PMPE expression plasmid M15 pREP (pQE-PmpE-Ct#37). The mature form of the *C. trachomatis* PMPE protein is expressed in *E. coli* as a fusion protein carrying the MRGS-(H)₆ domain encoded by the vector plasmid pOE-30 at the N-terminus.

Figure 2. Schematic map of the *C. trachomatis* PMPI expression plasmid TOP10 (pBAD-pmpI-Ct-Uni#7). The *C. trachomatis* PMPI protein is expressed in *E. coli* as a fusion protein carrying the HP-Thio domain encoded by the vector plasmid pBAD/Thio-E at the N-terminus.

Figure 3. A Coomassie blue-stained SDS-PAGE gel of the gel-purified *C*. 15 trachomatis PMPE protein expressed from the pQE-pmpE-Ct #37 plasmid. The PMPE protein migrates as an ~100 kDa protein. Pre-stained molecular weight markers (Lane 1) are Myosin (~250 kDa), Phosphorylase B (~148 kDa), BSA (~ 98 kDa), GDH (~64 kDa), ADH (~50 kDa), CAH (~36 kDA), Myoglobulin (30 kDa), Lysozyme (16 kDa), Aprotinin (~6 kDa), Insulin B chain (~4 kDa) (SeeBlue prestained standard Novex LC5625).

Figure 4. A Coomassie blue-stained SDS-PAGE gel of *E. coli* Top10 cell extracts carrying the *C. trachomatis* PMPI expression plasmid pBAD-pmpI-ct-Uni#7. Lane 1, pre-stained molecular weight markers (Novex MultiMark LC5725); lane 2, non-induced cells; lane 3, arabinose induced cells. The *C. trachomatis* PMPI protein in the arabinose-induced lane is indicated by an arrow. Molecular weight markers (Lane 1) are Myosin (~250 kDa), Phosphorylase B (~148 kDa), GDH (~60 kDa), CAH (~42 kDa), Myoglobulin-Blue (~30 kDa), Myoglobulin-Red (~22 kDa), Lysozyme(~17 kDa).

Figure 5. Full length nucleotide sequence and corresponding deduced amino acid sequence of the PMPE polypeptide of *Chlamydia trachomatis* L2.

Figure 6. Full length nucleotide sequence and corresponding deduced amino acid sequence of the PMPI polypeptide of *Chlamydia trachomatis* L2.

Figure 7. In vitro antigen-specific spleen cell proliferative response in animals immunized with recombinant PMPE and an adjuvant. Spleens were harvested from immunized C3H HeOuJ female mice approximately 14 days after a three dose immunization regimen. Single cell suspensions were prepared and the cells from two animals were pooled for analysis. Aliquots of the pooled cell samples were incubated for

72-96 hours in the presence of a test stimulant and pulsed with ³H-thymidine for the last 18-24 hours. Pooled samples incubated and pulsed labeled in parallel but in the absence of any stimulant served as the baseline ³H-thymidine uptake control. Represented is the stimulation index of cells stimulated with ConA (concanavalin A) (positive stipulation control); adjuvant (adjuvant employed in immunizations); PMPE (recombinant pmpE protein); EB (UV-inactivated *C. trachomatis* elementary bodies). 1 μg/ml (open bars), 4 μg/ml (solid bars) and 8 μg/ml (hatched bars) denote the three concentrations of *in vitro* stimulant used in the experiment. Stimulation index (SI) denotes the difference in ³H-thymidine incorporation of stimulated cells minus the background incorporation of the unstimulated controls. Bars denote the mean ± standard deviation in SI.

5. DETAILED DESCRIPTION OF THE INVENTION

5.1. CHLAMYDIA PMP POLYPEPTIDES

The present invention is generally directed to compositions and methods for the diagnosis, prevention, and treatment of Chlamydial infection. In one aspect, the composition of the subject invention provides pure native (wildtype) or recombinantly produced PMP polypeptides that comprise at least one immunogenic portion of a *Chlamydia* antigen.

The terms "treatment" or "therapy" as used herein and in the claims encompasses elimination as well as reduction in the severity or amelioration of disease symptoms caused directly or indirectly by the organism or numbers of organisms present.

In particular embodiments, the term "Chlamydia" refers to any Chlamydial species (spp.) including but not limited to Chlamydia trachomatis, Chlamydia pneumonia, Chlamydia psittaci and Chlamydia pecorum.

Strains from any of these organism may be obtained worldwide from any biologicals depository, particularly strains of *Chlamydia* ATCC VR-346, VR-347, VR-348B, VR-571B, VR-572, VR-573, VR-577, VR-578, VR-878, VF-879, VR-880, VR-885, VR-886, VR-887, VR-901B, VR-902B, VR-903, VR-1355, VR-1474, VR-1477, VR-2282, which may be obtained from the ATCC.

In a particular embodiment, the *Chlamydia* PMP protein is a PMPE polypeptide comprising or consisting of the amino acid sequence of SEQ ID NO.:2.

In another particular embodiment, the PMPE polypeptide is encoded by the nucleotide sequence of SEQ ID NO.:1. In another embodiment, the *Chlamydia* PMPE polypeptide comprises or consists of an amino acid sequence which is substantially

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homologous to SEQ ID NO.:2 or a portion thereof or is encoded by a nucleotide sequence substantially homologous to (or hybridizes under low, moderate or high stringency conditions to) the nucleotide sequence of SEQ ID NO.:1, or a portion thereof.

In another particular embodiment the *Chlamydia* PMP protein is a PMPI protein comprising or consisting of the amino acid sequence of SEQ ID NO.:4. In another embodiment, the *Chlamydia* polypeptide is a PMPI polypeptide encoded by the nucleotide sequence of SEQ ID NO.:3. In another embodiment, the *Chlamydia* polypeptide comprises or consists of an amino acid sequence which is substantially homologous to SEQ ID NO.:4, or a portion thereof, or is encoded by a nucleotide sequence substantially homologous to (or hybridizes under low, moderate or high stringency conditions to) the nucleotide sequence of SEQ ID NO.:3, or a portion thereof.

The present invention provides the family of *Chlamydia* PMPE and PMPI proteins. The amino acid sequences of SEQ ID NOs.:2 and 4 and the nucleotide sequences of SEQ ID NOs.:1 and 3 represent the amino acid and nucleotide sequences, respectively, of the PMPE and PMPI polypeptides, respectively, of the *Chlamydia trachomatis* L2 serovar. The invention also relates to PMPE and PMPI polypeptide homologs from other *Chlamydia trachomatis* serovars and other *Chlamydia* species, as well as other derivatives, analogs, and fragments thereof (*i.e.*, PMPE- and PMPI-derived polypeptides). In a preferred embodiment, immunization with the polypeptide of the invention elicits antibodies that specifically bind to PMPE or PMPI polypeptides from other *Chlamydia* serovars and, preferably, species (preferably all species, but may be a subset of species) besides the serovar and species from which the polypeptide was isolated or derived.

As used herein a "substantially homologous" sequence is at least 70%, preferably greater than 80%, more preferably greater than 90% identical to a reference amino acid or nucleotide sequence of identical size or when compared to a reference sequence when the alignment or comparison is conducted by a computer homology program or search algorithm known in the art. By way of example and not limitation, useful computer homology programs include the following: Basic Local Alignment Search Tool (BLAST) (www.ncbi.nlm.nih.gov) (Altschul et al., 1990, J. of Molec. Biol., 215:403-410, "The BLAST Algorithm"; Altschul et al., 1997, Nuc. Acids Res. 25:3389-3402) a heuristic search algorithm tailored to searching for sequence similarity which ascribes significance using the statistical methods of Karlin and Altschul 1990, Proc. Nat'l Acad. Sci. USA, 87:2264-68; 1993, Proc. Nat'l Acad. Sci. USA 90:5873-77. Five specific BLAST programs perform the following tasks:

- 1) The BLASTP program compares an amino acid query sequence against a protein sequence database.
- 2) The BLASTN program compares a nucleotide query sequence against a nucleotide sequence database.
- 5 3) The BLASTX program compares the six-frame conceptual translation products of a nucleotide query sequence (both strands) against a protein sequence database.
 - 4) The TBLASTN program compares a protein query sequence against a nucleotide sequence database translated in all six reading frames (both strands).
- 5) The TBLASTX program compares the six-frame translations of a nucleotide query sequence against the six-frame translations of a nucleotide sequence database.

Smith-Waterman (database: European Bioinformatics Institute wwwz.ebi.ac.uk/bic_sw/) (Smith-Waterman, 1981, <u>J. of Molec. Biol.</u>, 147:195-197) is a mathematically rigorous algorithm for sequence alignments.

FASTA (see Pearson et al., 1988, <u>Proc. Nat'l Acad. Sci. USA</u>, 85:2444-2448)

15 is a heuristic approximation to the Smith-Waterman algorithm. For a general discussion of the procedure and benefits of the BLAST, Smith-Waterman and FASTA algorithms see Nicholas et al., 1998, "A Tutorial on Searching Sequence Databases and Sequence Scoring Methods" (www.psc.edu) and references cited therein.

By further way of example and not limitation, useful computer homology 20 algorithms and parameters for determining percent identity include the following:

To determine the percent identity of two amino acid sequences or of two nucleotide sequences, *e.g.*, between PMP sequences and other known sequences, the sequences are aligned for optimal comparison purposes (*e.g.*, gaps can be introduced in the sequence of a first amino acid or nucleic acid sequence for optimal alignment with a second amino or nucleic acid sequence). The amino acid residues or nucleotides at corresponding amino acid positions or nucleotide positions are then compared. When a position in the first sequence is occupied by the same amino acid residue or nucleotide as the corresponding position in the second sequence, then the molecules are identical at that position. The percent identity between the two sequences is a function of the number of identical positions shared by the sequences (*i.e.*, % identity = # of identical positions/total # of positions (*e.g.*, overlapping positions) x 100). In one embodiment, the two sequences are the same length.

The determination of percent identity between two sequences can be accomplished using a mathematical algorithm, a preferred, non-limiting example of a mathematical algorithm utilized for the comparison of two sequences is the algorithm of

Karlin and Altschul, 1990, <u>Proc. Nat'l Acad. Sci. USA</u>, 87:2264-68; as modified by 1993, <u>Proc. Nat'l Acad. Sci. USA</u> 90:5873-77. Such algorithm is incorporated into the NBLAST and XBLAST programs of Altschul, 1990, <u>J. of Molec. Biol.</u>, 215:403-410. BLAST nucleotide searches can be performed with the NBLAST program, score = 100, wordlength

- = 12 to obtain nucleotide sequences homologous to a nucleic acid molecule of the invention. BLAST protein searches can be performed with the XBLAST program, score = 50, wordlength = 3 to obtain amino acid sequences homologous to a protein molecule of the invention. To obtain gapped alignments for comparison purposes, Gapped BLAST can be utilized as described in Altschul, 1997, Nuc. Acids Res. 25:3389-3402. Alternatively, PSI-
- 10 BLAST can be used to perform an iterated search which detects distant relationships between molecules (*Id.*). When utilizing BLAST, Gapped BLAST, and PSI-BLAST programs, the default parameters of the respective programs can be used. Another preferred, non-limiting example of a mathematical algorithm utilized for the comparison of sequences is the algorithm of Myers and Miller, CABIOS (1989). Such an algorithm is
- incorporated into the ALIGN program (version 2.0) which is part of the CGC sequence alignment software package. When using the ALIGN program for comparing amino acid sequences, a PAM120 weight residue table, a gap length penalty of 12, and a gap penalty of 4 can be used. Additional algorithms for sequence analysis are known in the art and include ADVANCE and ADAM as described in Torellis and Robotti (1994) Comput. Appl. Biosc.,
- 20 10:3-5; and FASTA described in Pearson and Lipman, 1988, Proc. Nat'l Acad. Sci. USA, 85:2444-2448. Within FASTA, ktup is a control option that sets the sensitivity and speed of the search. If ktup = 2, similar regions in the two sequences being compared are found by looking at pairs of aligned residues; if ktup = 1, single aligned amino acids are examined. Ktup can be set to 2 or 1 for protein sequences, or from 1 to 6 for nucleotide sequences.
- The default, if ktup is not specified, is 2 for proteins and 6 for nucleotides. For a further description of FASTA parameters, see http://bioweb.pasteur.fr/docs/man/man/fasta.1.html#sect2, the contents of which are incorporated herein by reference. Alternatively, protein sequence alignment may be carried out using the CLUSTAL W algorithm as described by Higgins et al., 1996, Methods
- 30 Enzymol., 266:383-402.

The percent identity between two sequences can be determined using techniques similar to those described above, with or without allowing gaps. In calculating percent identity, only exact matches are counted.

According to various aspects of the invention, the polypeptides of the invention are characterized by their apparent molecular weights based on the polypeptides'

migration in SDS-PAGE (sodium dodecyl sulfate-polyacrylamide gel electrophoresis) relative to the migration of known molecular weight markers. While any molecular weight standards known in the art may be used with the SDS-PAGE, preferred molecular weight markers comprise pre-stained Myosin (250 kDa), Phosphorylase B (148 kDa), BSA (98 kDa) and GDH (64 kDa). One skilled in the art will appreciate that the polypeptides of the invention may migrate differently in different types of gel systems (e.g., different buffers; different types and concentrations of gel, crosslinkers or SDS, etc.). One skilled in the art will also appreciate that the polypeptides may have different apparent molecular weights due to different molecular weight markers used with the SDS-PAGE. Hence, the molecular weight characterization of the polypeptides of the invention is intended to be directed to cover the same polypeptides on any SDS-PAGE system and with any set of molecular weight markers which might indicate slightly different apparent molecular weights for the polypeptides than those disclosed herein.

In specific embodiments, the subject invention discloses PMP polypeptides comprising an immunogenic portion of a *Chlamydia* antigen, wherein the *Chlamydia* antigen comprises an amino acid sequence encoded by a nucleic acid molecule comprising a sequence selected from the group consisting of (a) nucleotide sequences of SEQ ID NO.:1 or SEQ ID NO.:3, (b) the complements of said nucleotide sequences; and (c) variants of such sequences.

Nucleic acid molecules encoding the precursor and mature forms of *Chlamydia* polypeptides are provided by the invention.

5.2 CHLAMYDIA PMP DERIVED POLYPEPTIDES

The term "antigens" and its related term "antigenic" as used herein and in the claims refers to a substance to which an antibody or T-cell receptor specifically binds. As used herein, antisera, antibodies and T-cells are "antigen-specific" if they specifically bind to or react with an antigen and do not react detectably with unrelated proteins other than by non-specific interaction. Preferably said antigens are immunogenic.

The term "immunogenic" as used herein and in the claims refers to the ability to induce an immune response, *e.g.*, an antibody and/or a cellular immune response in an animal, preferably a mammal or a bird.

In a specific embodiment of the invention, proteins consisting of or comprising a fragment of a PMPE protein consisting of at least 8 (continuous) amino acids of the protein are provided. In other embodiments, the fragment consists of at least 10, 20, 40, 50, 60, 80, 100, 150, 200, 300, 400 or 500 amino acids of SEQ ID NO.:2. In specific

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embodiments, such fragments are not larger than 10, 11, 12, 15, 20, 25, 35, 50, 75, 80, 90, 100, 125, 150, 175, 200, 225, 250, 275, 300, 325, 350, 400, 425, 450, 475, 500, 525, 550, 575, 600, 625, 650, 675 or 700 amino acids. In preferred embodiments, the PMPE-derived polypeptide contains a sequence forming an outer-surface epitope.

In a particular embodiment, the PMPE-derived polypeptide is a fragment of a PMPE polypeptide which fragment comprises or consists of an amino acid sequence of any of SEQ ID NOs.:5-22. In another particular embodiment, the PMPE-derived polypeptide is a fragment of a PMPE polypeptide, which fragment comprises or consists of an amino acid sequence of any of SEQ ID NOs.:5-22, but also comprises additional C-terminal or N-10 terminal PMPE sequences.

In a specific embodiment of the invention, proteins are provided that consist of or comprise a fragment of a PMPI protein consisting of at least 8 (continuous) amino acids of SEQ ID NO.:4. In other embodiments, the fragment consists of at least 10, 15, 20, 25, 50, 75, 100, 150 or 200 amino acids of SEQ ID NO.:4. In specific embodiments, such 15 fragments are not longer than 10, 11, 12, 15, 20, 25, 35, 50, 75, 80, 90, 100, 125, 150, 175, 200, 225, 250, 275, 300, 325, 350, 400, 425, 450, 475, 500, 525, 550, 575, 600, 625, 650, 675 or 700 amino acids. In preferred embodiments, PMPI-derived polypeptide contains a sequence forming an outer-surface epitope.

In a particular embodiment, the PMPI-derived polypeptide is a fragment of a 20 PMPI peptide which comprises the amino acid sequence of any of SEQ ID NOs.:23-34.

In another particular embodiment, the PMPE-derived polypeptide is a fragment of a PMPE polypeptide which comprises the amino acid sequence of any of SEQ ID NOs.:23-34, but also comprises additional C-terminal or N-terminal PMPI sequences.

Preferably, the PMPE-derived polypeptides of the invention are

25 immunologically cross-reactive with the PMPE polypeptide, and are capable of eliciting in an animal an immune response to Chlamydia, Chlamydia elemental bodies (EB), Chlamydia reticulate bodies (RBs), Chlamydia infected cells or antigen presenting cells expressing Chlamydial antigens and/or are able to be bound by anti-PMPE antibodies.

Preferably, the PMPI-derived polypeptides of the invention are 30 immunologically cross-reactive with the PMPI polypeptide, and are capable of eliciting in an animal an immune response to Chlamydia, Chlamydia elemental bodies (EB), reticulocyte bodies (RBs), Chlamydia infected cells or antigen presenting cells expressing Chlamydial antigens and/or are able to be bound by anti-PMPI antibodies. More preferably, the PMP-derived polypeptides of the invention comprise sequences forming one or more

35 epitopes of the native PMPE or PMPI polypeptide of Chlamydia (i.e., the epitopes of PMPE

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Polypeptide derivatives can also be constructed by deletions that remove a part of the parent polypeptide, while retaining the desired specific antigenicity. Deletions can also remove regions of high variability among strains.

Also preferably, the Chlamydia derived polypeptides of the invention 15 comprise sequences that form one or more epitopes of a native Chlamydia polypeptide that elicit bactericidal or opsonizing antibodies. Such preferred Chlamydia-derived polypeptides may be identified by their ability to generate antibodies that kill Chlamydia spp., particularly, Chlamydia trachomatis cells. For example, polypeptides from a limited 20 or complete protease digestion or chemical cleavage of a Chlamydia polypeptide are fractionated using standard methods, (e.g., by limited proteolytic digestion using enzymes such as trypsin, papain, or related proteolytic enzymes or by chemical cleavage using agents such as cyanogen bromide and followed by fractionation of the digestion or cleavage products), injected into animals and the antibodies produced therefrom are tested for the 25 ability to interfere with or kill Chlamydia cells and/or Chlamydia infected cells. Once identified and isolated, the amino acid sequences of such preferred Chlamydia-derived polypeptides are determined using standard sequencing methods. The determined sequence may be used to enable production of such polypeptides by synthetic chemical and/or genetic engineering means.

These preferred *Chlamydia*-derived polypeptides also can be identified by using anti-whole cell antibodies to screen bacterial libraries expressing random fragments of *Chlamydia* genomic DNA or cloned nucleotide sequences encoding a PMPE or PMPI polypeptide or fragments thereof. See, *e.g.*, Sambrook et al., 1989, Molecular Cloning, A Laboratory Manual, 2nd ed., Cold Spring Harbor Press, Cold Spring Harbor, New York, Vol. 1, Chapter 12. The reactive clones are identified and their inserts are isolated and

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sequenced to determine the amino acid sequences of such preferred *Chlamydia*-derived polypeptides.

Examples of immunogenic portions of antigens contemplated by the present invention include polypeptides comprising or consisting of the fragments set forth in Tables 1 and 2, where the numbers following the PMPE (table 1, column 1) or PMPI (table 2, column 1) designation refer to the amino acid residues in SEQ ID NOs.:2 or 4, respectively. Polypeptides comprising at least an immunogenic portion of one or more *Chlamydia* antigens or immunogenic portions as described herein may generally be used, alone or in combination, to detect, prevent, treat or reduce the severity of Chlamydial infection.

TABLE 1: PREFERRED FRAGMENTS OF PMPE

	FRAGMENT	SEQ ID NO.:
15	PMPE15-56 PMPE15-121 PMPE45-125 PMPE125-190 PMPE195-261	5 6 7 8 9
20	PMPE275-366 PMPE375-440 PMPE440-490 PMPE525-590 PMPE590-625 PMPE615-650	10 11 12 13 14 15
25	PMPE625-700 PMPE725-800 PMPE755-775 PMPE785-845 PMPE815-865 PMPE1-31 PMPE1-500	16 17 18 19 20 21 22

30

35

TABLE 2: PREFERRED FRAGMENTS OF PMPI

	<u>FRAGMENT</u>	SEQ ID NO.:
	PMPI 13-40	23
5	PMPI 70-110	24
	PMPI 150-225	25
	PMPI 250-290	26
	PMPI 370-455	27
	PMPI 400-455	28
	PMPI 470-520	29
1.0	PMPI 615-670	30
10	PMPI 710-775	31
	PMPI 765-825	32
	PMPI 830-860	33
	PMP 1-500	34

Polypeptides having a sequence homologous to one of the PMP polypeptides of the invention, including naturally-occurring allelic variants, as well as mutants, variants or any other non-naturally occurring variants that are analogous to, *e.g.*, cross-reacting with antibodies against, a PMP polypeptide of the present invention, are encompassed by the present invention.

Allelic variants are very common in nature. For example, a bacterial species 20 e.g., C. trachomatis, is usually represented by a variety of strains or serovars that differ from each other by minor allelic variations. Indeed, a polypeptide that fulfills the same biological function in different strains can have an amino acid sequence that is not identical in each of the strains. Such an allelic variation may be equally reflected at the nucleic acid molecule level.

An allelic variant is an alternate form of a polypeptide that is characterized as having a substitution, deletion, or addition of one or more amino acids that does not substantially alter the biological function of the polypeptide. By "biological function" is meant the function of the polypeptide in the cells in which it naturally occurs, even if the function is not necessary for the growth or survival of the cells.

Nucleic acid molecules, *e.g.*, DNA molecules, encoding allelic variants can easily be retrieved by the polymerase chain reaction (PCR) amplification of genomic bacterial DNA extracted by conventional methods. This involves the use of synthetic oligonucleotide primers matching upstream and downstream sequences of the 5' and 3' ends of the encoding domains. Typically, a primer can consist of 10 to 40, preferably 15 to

35 25 nucleotides. It may be also advantageous to select primers containing C and G

nucleotides in a proportion sufficient to ensure efficient hybridization; e.g., an amount of C and G nucleotides of at least 40%, preferably 50% of the total number of nucleotides in the primer.

Variants of Chlamydia trachomatis PMP proteins, such as PMP proteins from the A, B, Ba, C, D, Da, E, F, G, H, I, Ia, J, K, MoPN, L1, L2, and L3 serovars, which share sequence homology to the PMP polypeptides and nucleic acid molecules described herein are also provided.

Homology is defined as being at least 70, 80, 85, 90, 95 or 99% identical to a reference sequence of identical size or when the alignment or comparison is by a computer 10 homology program or search algorithm known in the art (see Section 5.1 supra). Preferably, the serovar homologues show 70, 80, 85, 90, 95 or 99% homology to the corresponding polypeptide sequence(s) described herein. Most preferably, the serovar homologues show 95-99% homology to the corresponding polypeptide sequence(s) described herein.

A Chlamydia-derived PMP polypeptide of the invention may also be a 15 modified PMPE or PMPI polypeptide or fragment thereof (i.e., a Chlamydia polypeptide or fragment having one or more amino acid substitutions, insertions and/or deletions of the wild-type Chlamydia sequence or amino acids chemically modified in vivo or in vitro). Such modifications may enhance the immunogenicity of the resultant Chlamydia-derived 20 polypeptide product or have no effect on such activity.

As used herein, the term "enhance the immunogenicity" refers to an increased antibody titer or increased cellular immune response elicited by exposure to the modified polypeptide as compared to the immune response elicited by unmodified polypeptides or formalin or glutaraldehyde fixed Chlamydia. Modification techniques that 25 may be used include, but are not limited to, those disclosed in U.S. Patent No. 4,526,716.

As an illustrative, non-limiting example, one or more amino acid residues within the PMP-derived polypeptide sequence can be substituted by another amino acid of a similar polarity which acts as a functional equivalent, resulting, for example, in a silent alteration. Substitutes for an amino acid within the sequence may be selected from other 30 members of the class to which the amino acid belongs. For example, the nonpolar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan and methionine. The polar neutral amino acids include glycine, serine, threonine, cysteine, tyrosine, asparagine, and glutamine. The positively charged (basic) amino acids include arginine, lysine and histidine. The negatively charged (acidic)

35 amino acids include aspartic acid and glutamic acid.

Included within the scope of the invention are PMP-derived polypeptides which are Chlamydia PMP polypeptide fragments or other derivatives or analogs which are differentially modified during or after translation, e.g., by glycosylation, acetylation, phosphorylation, lipidation, amidation, derivatization by known protecting/blocking groups, proteolytic cleavage, linkage to an antibody molecule or other cellular ligand, etc. Any of numerous chemical modifications may be carried out by known techniques, including but not limited to, specific chemical cleavage by cyanogen bromide, trypsin, chymotrypsin, papain, V8 protease, NaBH₄, acetylation, formylation, oxidation, reduction, metabolic synthesis in the presence of tunicamycin, etc.

Furthermore, if desired, nonclassical amino acids or chemical amino acid analogs can be introduced as a substitution or addition into the Chlamydia polypeptide sequence. Non-classical amino acids include but are not limited to the D-isomers of the common amino acids, α-amino isobutyric acid, 4-aminobutyric acid, Abu, 2-amino butyric acid, γ-Abu, ε-Ahx, 6-amino hexanoic acid, Aib, 2-amino isobutyric acid, 3-amino 15 propionic acid, ornithine, norleucine, norvaline, hydroxyproline, sarcosine, citrulline, cysteic acid, t-butylglycine, t-butylalanine, phenylglycine, cyclohexylalanine, \(\beta-alanine, fluoro-amino acids, designer amino acids such as β-methyl amino acids, Cα-methyl amino acids, Nα-methyl amino acids, PNA's and amino acid analogs in general. Furthermore, the amino acid can be D (dextrorotary) or L (levorotary).

20 A Chlamydia PMP-derived polypeptide may further be a chimeric polypeptide comprising one or more heterologous polypeptides, lipids, phospholipids or lipopolysaccharides of Chlamydia origin or of another bacterial or viral origin, fused (e.g., covalently bound) to the amino-terminus or carboxyl-terminus or is within a complete PMPE or PMPI polypeptide or PMP-derived polypeptide. Useful heterologous 25 polypeptides to be included within such a chimeric polypeptide include, but are not limited to, a) pre- and/or pro- sequences that facilitate the transport, translocation and/or processing of the PMP-derived polypeptide in a host cell, b) affinity purification sequences, and c) any useful immunogenic sequences (e.g., sequences encoding one or more epitopes of a surfaceexposed protein of a microbial pathogen). One preferred heterologous protein of the 30 chimeric polypeptide includes Hin47 (see U.S. Patents 5,679,547 and 5,721,115 which are hereby incorporated by reference in their entirety). Another preferred chimeric polypeptide includes an HMWP protein of Chlamydia or fragments thereof (see United States patent application Serial No. 08/942,596, filed October 2, 1997, entitled "Chlamydia Protein, Gene Sequence and Uses Thereof", which is incorporated by reference herein in its entirety) or

Chlamydia MOMP protein or fragments thereof (see U.S. Patent No. 5,869,608, which is incorporated herein by reference in its entirety).

PMP-derived polypeptides also include but are not limited to fusion polypeptides comprising at least two regions derived from one or more Chlamydia proteins, each having T-cell or antibody stimulating activity. The regions may be derived from the same Chlamydia protein or may comprise one or more regions from more than one Chlamydia protein. The polypeptides are arranged in a nonsequential order or noncontiguous order (e.g., in an order different from the order of the amino acids of the native protein). A preferred polypeptide of the invention is a fusion polypeptide comprising 10 at least two peptides, each of which peptides consists of an amino acid sequence selected from the group consisting of SEQ ID NOs.:5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, and 34, with the proviso that the peptides are arranged in a configuration that is different from the configuration of a naturally occurring PMPE or PMPI polypeptide.

Other preferred PMP-derived polypeptides of the invention are an isolated fusion polypeptide wherein the polypeptide comprises all the peptides consisting of the amino acid sequences of SEQ ID NOs.:5, 6, 7, 8, 9, 10 and 11, or an isolated fusion polypeptide wherein the polypeptide comprises all the peptides consisting of the amino acid sequences of SEQ ID NOs.:23, 24, 25, 26, 27, 28, and 29, with the proviso that the peptides 20 are arranged in a configuration that is different from the configuration of a naturally occurring PMPE or PMPI polypeptide. A preferred PMP-derived polypeptide is an isolated fusion polypeptide comprising a peptide consisting of an amino acid sequence of any of SEQ ID NOs.:5, 6, 7, 8, 9, 10, 11, 23, 24, 25, 26, 27, 28 or 29, with the proviso that the peptides are arranged in a configuration that is different from the configuration of a 25 naturally occurring PMPE or PMPI polypeptide.

If desired, the amino acid sequences of the regions can be produced and joined by a linker.

Suitable peptide linker sequences may be chosen based on the following factors: (1) their ability to adopt a flexible extended conformation; (2) their ability to adopt 30 a secondary structure that could interact with functional epitopes of the first and second polypeptides; (3) the lack of hydrophobic or charged residues that might react with the polypeptide functional epitopes; (4) the ability to increase solubility; and (5) the ability to increase sensitivity to processing by antigen-presenting cells. Such linkers can be any amino acid sequence or other appropriate link or joining agent.

35

Linkers useful in the invention include linkers comprising a charged amino acid pair such as KK or RR, linkers sensitive to cathepsin and/or other trypsin-like enzymes, thrombin or Factor X_{a} , or linkers which result in an increase in solubility of the polypeptide.

Preferred peptide linker sequences contain Gly, Asn and Ser residues.

Amino acid sequences which may be usefully employed as linkers include those disclosed in Maratea et al. Gene 40:39-46 (1985); Murphy et al., Proc. Nat. Acad Sci USA 83:8258-8562 (1986); U.S. Patent No. 4,935,233 and U.S. Patent No. 4,751,180. The linker sequence may be from 1 to about 50 amino acids in length.

Another particular example of fusion polypeptides of the invention includes a polypeptide or polypeptide derivative of the invention fused to a polypeptide having adjuvant activity, such as the subunit B of either cholera toxin or *E. coli* heat labile toxin. Another particular example of a fusion polypeptide encompassed by the invention includes a polypeptide or polypeptide derivative of the invention fused to a cytokine (such as, but not limited to, IL-2, IL-4, IL-10, IL-12, or interferon). A polypeptide of the invention can be fused to the N- or C-terminal end of the polypeptide having adjuvant activity.

Alternatively, a polypeptide of the invention can be fused within the amino acid sequence of the polypeptide having adjuvant activity.

Also preferably, the *Chlamydia* derived fusion polypeptides of the invention comprise sequences that form one or more epitopes of a native *Chlamydia* polypeptide that elicit bactericidal or opsonizing antibodies and/or T-cells. Such preferred *Chlamydia*-derived polypeptides may be identified by their ability to generate antibodies and/or T-cells that kill cells infected with *Chlamydia* spp. cells particularly, *Chlamydia trachomatis* cells.

5.3 <u>ISOLATION AND PURIFICATION OF PMP POLYPEPTIDES AND PMP-DERIVED POLYPEPTIDES</u>

The invention provides isolated PMPE and PMPI polypeptides, PMPE-derived and PMPI-derived polypeptides. As used herein, the term "isolated" means that the product is significantly free of other biological materials with which it is naturally associated, or free from other biological materials derived, for example, from a recombinant host cell that has been genetically engineered to express the polypeptide of the invention. That is, for example, an isolated PMP polypeptide composition that is between about 70% and 99% pure PMP polypeptide by weight. As used herein, the term "purified" means that the product is substantially free of other biological material with which it is naturally associated, or free from other biological materials derived, for example, from a recombinant host cell that has been genetically engineered to express the polypeptide of the invention.

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That is, a purified PMP polypeptide composition is at least 70-95% pure PMP polypeptide by weight, preferably at least 98% pure PMP polypeptide by weight, and most preferably at least 99% pure PMP polypeptide by weight.

The PMP polypeptide of the invention may be isolated from a protein extract, including a whole cell extract of any Chlamydia spp., including, but not limited to, Chlamydia trachomatis, Chlamydia pneumoniae, Chlamydia pecorum, and Chlamydia psittaci. Strains from any of these organisms may be obtained worldwide from any biologicals depository, for example, strains of ATCC VR-346, VR-347, VR-348B, VR-571B, VR-572, VR-573, VR-577, VR-578, VR-878, VF-879, VR-880, VR-885, VR-886, 10 VR-887, VR-901B, VR-902B, VR-903, VR-1355, VR-1474, VR-1477, or VR-2282 may be obtained from the American Type Culture Collection.

Another source of the PMP polypeptide is a protein preparation from a gene expression system (such as E. coli) engineered to express a cloned sequence encoding a PMP polypeptide or PMP-derived polypeptide (see Section 5.5, *infra*).

The PMP polypeptide can be isolated and purified from the source material using any biochemical technique and approach well known to those skilled in the art. In one approach, Chlamydia cellular envelope is obtained by standard techniques and inner membrane, periplasmic and outer membrane proteins are solubilized using a solubilizing compound such as a detergent or hypotonic solution. A preferred detergent solution is one 20 containing octyl glucopyranoside (OG), sarkosyl or TRITON X100™. A preferred solubilizing hypotonic solution is one containing LiCl. The PMP polypeptide is in the solubilized fraction. Cellular debris and insoluble material in the extract are separated and removed preferably by centrifugation. The polypeptides in the extract are concentrated, incubated in SDS-containing Laemmli gel sample buffer at 100°C for 5 minutes and then 25 fractionated by electrophoresis in a denaturing sodium dodecylsulfate (SDS) polyacrylamide gel from about 6% to about 12%, with or without a reducing agent. See Laemmli, 1970, Nature 227:680-685. The band or fraction identified as a PMP polypeptide, having an apparent molecular weight of about 90-115 kDa, as described above, may then be isolated directly from the fraction or gel slice containing the PMP polypeptide. In a preferred 30 embodiment, the PMP polypeptide has an apparent molecular weight of about 90-115 kDa which can be determined by comparing its migration distance or rate in denaturing SDS-PAGE relative to the migration of known molecular weight markers such as of myosin (250 kDa), Phosphorylase B (148 kDa), BSA (98 kDa) and GDH (64 kDa) (weights for prestained markers).

Another method of purifying PMP polypeptide is by affinity chromatography using anti-PMP antibodies (see Section 5.4). The affinity chromatography may be carried out using either polyclonal or monoclonal anti-PMP antibodies, preferably, monoclonal antibodies. The antibodies are covalently linked to agarose gels activated by cyanogen bromide or succinamide esters (Affi-Gel, BioRad, Inc.) or by other methods known to those skilled in the art. The protein extract is loaded on the top of the gel and is left in contact with the gel for a period of time and under standard reaction conditions sufficient for PMP polypeptide to bind to the antibody. Preferably, the solid support is a material used in a chromatographic column. The affinity gel is washed to remove other proteins and cell materials not bound by the anti-PMP antibody. The PMP polypeptide is then removed from the antibody to recover the PMP polypeptide in isolated, or preferably, in purified form.

A PMP-derived polypeptide of the invention can be produced by chemical and/or enzymatic cleavage or degradation of an isolated or purified PMP polypeptide. A PMP-derived polypeptide can also be chemically synthesized based on the known amino acid sequence of the PMP polypeptide and, in the case of a chimeric polypeptide, the amino acid sequence of the heterologous polypeptide, by methods well known in the art. See, for example, Creighton, 1983, Proteins: Structures and Molecular Principles, W.H. Freeman and Co., NY.

A PMP-derived polypeptide can also be produced in a gene expression system expressing a recombinant nucleic acid construct comprising a sequence encoding a PMP-derived polypeptide. The nucleotide sequences encoding polypeptides of the invention may be synthesized, and/or cloned, and expressed according to techniques well known to those skilled in the art. See, for example, Sambrook, et al., 1989, Molecular Cloning, A Laboratory Manual, Vols. 1-3, Cold Spring Harbor Press, NY, Chapter 9.

25 PMP-derived polypeptides of the invention can be fractionated and purified by the application of standard protein purification techniques, modified and applied in accordance with the discoveries and teachings described herein. In particular, preferred PMP polypeptides of the invention, those that form an outer-surface or exposed epitope of the native PMP polypeptide, may be isolated and purified according to the affinity procedures disclosed above for the isolation and purification of a PMP polypeptide (*e.g.*, affinity purification using anti-PMP antibodies).

If desirable, the polypeptides of the invention may be further purified using standard protein or peptide purification techniques including but not limited to, electrophoresis, centrifugation, gel filtration, precipitation, dialysis, chromatography

35 (including ion exchange chromatography, affinity chromatography, immunoadsorbent

affinity chromatography, dye-binding chromatography, size exclusion chromatography, hydroxyappitite chromatography, reverse-phase high performance liquid chromatography, and gel permeation high performance liquid chromatography), isoelectric focusing, and variations and combinations thereof.

One or more of these techniques may be employed sequentially in a procedure designed to isolate and/or purify the PMP polypeptide or the PMP-derived polypeptides of the invention according to its/their physical or chemical characteristics. These characteristics include the hydrophobicity, charge, binding capability, and molecular weight of the protein. The various fractions of materials obtained after each technique are 10 tested for binding to the PMP receptor or ligand or to anti-PMP antibodies or for functional activity ("test" activities). Those fractions showing such test activity are then pooled and subjected to the next technique in the sequential procedure, and the new fractions are tested again. The process is repeated until fractions are obtained that have one or more of the above described "test" activities and that contain only a single band (or a very predominant 15 band) or entity when subjected to polyacrylamide gel electrophoresis or chromatography.

5.4. PMP IMMUNOGENS AND ANTI-PMP ANTIBODIES

The present invention provides antibodies that specifically bind a PMP polypeptide and/or PMP-derived polypeptide. For the production of such antibodies, 20 isolated or, preferably, purified preparations of a PMP polypeptide or PMP-derived polypeptide are used as immunogens in an immunogenic composition.

In one embodiment, the PMP polypeptide is separated from other outer membrane or periplasmic proteins present in the extracts of Chlamydia cells using SDS-PAGE (see Section 5.3 above) and the gel slice containing the PMP polypeptide is used as 25 an immunogen and injected into a rabbit to produce antisera containing polyclonal PMP antibodies. The same immunogen can be used to immunize mice for the production of hybridoma lines that produce monoclonal anti-PMP antibodies. In particular embodiments, the immunogen is an SDS-PAGE gel slice containing isolated or purified PMP from any Chlamydia strain, including, but not limited to, Chlamydia trachomatis, Chlamydia 30 pneumoniae, Chlamydia pecorum, and Chlamydia psittaci. Particularly preferred are the strains of Chlamydia trachomatis ATCC: VR-346, VR-347, VR-348B, VR-571B, VR-572, VR-573, VR-577, VR-578, VR-878, VF-879, VR-880, VR-885, VR-886, VR-887, VR-901B, VR-902B, VR-903, VR-1355, VR-1474, VR-1477, VR-2282.

In other embodiments, peptide fragments of a PMP polypeptide are used as 35 immunogens. Preferably, peptide fragments of purified PMP polypeptide are used. The

peptides may be produced by protease digestion, chemical cleavage of isolated or purified PMP polypeptide, chemical synthesis or by recombinant expression, after which they are then isolated or purified. Such isolated or purified peptides can be used directly as immunogens. In particular embodiments, useful peptide fragments are 8 or more amino acids in length.

Useful immunogens may also comprise such peptides or peptide fragments conjugated to a carrier molecule, preferably a carrier protein. Carrier proteins may be any commonly used in immunology, include, but are not limited to, bovine serum albumin (BSA), chicken albumin, keyhole limpet hemocyanin (KLH), tetanus toxoid and the like.

10 For a discussion of hapten protein conjugates, see, for example, Harlow and Lane

Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory Press, Cold Spring
Harbor, NY, 1988, or a standard immunology textbook such as Roitt, et al.,

IMMUNOLOGY, C.V. Mosby Co., St. Louis, MO (1985) or Klein, IMMUNOLOGY,
Blackwell Scientific Publications, Inc., Cambridge, MA (1990).

In yet another embodiment, for the production of antibodies that specifically bind one or more epitopes of a native PMP polypeptide, intact *Chlamydia* cells or elemental bodies (EBs) or reticulate bodies (RBs) prepared therefrom, or cells infected with *Chlamydia* are used as immunogen. The cells, EBs, RBs or cells infected with *Chlamydia* may be fixed with agents such as formaldehyde or glutaraldehyde before immunization.

20 See Harlow and Lane, <u>Antibodies: A Laboratory Manual</u>, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1988, Chapter 15. It is preferred that such anti-whole cell antibodies be monoclonal antibodies. Hybridoma lines producing the desired monoclonal antibodies can be identified by using purified PMP polypeptide, intact *Chlamydia* cells, EBs, RBs or cells infected with *Chlamydia* as the screening ligand. The immunogen for

25 inducing these antibodies may be whole *Chlamydia* cells, EBs, RBs, extracts or lysates of any *Chlamydia*, including, but not limited to, *Chlamydia trachomatis*, *Chlamydia pneumoniae*, *Chlamydia pecorum*, and *Chlamydia psittaci*. Particularly preferred are strains of *Chlamydia* strains ATCC VR-346, VR-347. VR-348B, VR-571B, VR-572, VR-573, VR-577, VR-578, VR-878, VF-879, VR-880, VR-885, VR-886, VR-887, VR-901B, VR-30 902B, VR-903, VR-1355, VR-1474, VR-1477, VR-2282.

Polyclonal antisera produced by immunization with cells infected with *Chlamydia*, whole cells, EBs or RBs contain antibodies that bind other *Chlamydia* proteins ("non-anti-PMP antibodies") and thus are more cumbersome to use when it is known or suspected that the sample contains other *Chlamydia* proteins or materials that are cross-

35 reactive with these other proteins. Under such circumstances, any binding by the anti-whole

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cell antibodies of a given sample or band must be verified by coincidental binding of the same sample or band by antibodies that specifically bind the PMP polypeptide (e.g., anti-PMPE antibodies) and/or a PMP-derived polypeptide, or by competition tests using anti-PMP antibodies, PMP polypeptides or PMP-derived polypeptides as the competitor (i.e., addition of anti-PMP antibodies, PMP polypeptide or PMP-derived polypeptide to the reaction mix lowers or abolishes sample binding by anti-whole cell antibodies). Alternatively, such polyclonal antisera containing "non-anti-PMP" antibodies, may be cleared of such non-anti-PMP antibodies by standard approaches and methods. For example, the non-anti-PMP antibodies may be removed by precipitation with cells having a 10 deletion of the PMP coding sequence or Chlamydia strains known not to have the PMP polypeptide; or by absorption to columns comprising such cells or outer membrane proteins of such cells.

In further embodiments, useful immunogens for eliciting antibodies of the invention comprise mixtures of two or more of any of the above-mentioned individual 15 immunogens.

Immunization of animals with the immunogens described herein, preferably of humans, rabbits, rats, ferrets, mice, sheep, goats, cows or horses, is performed following procedures well known to those skilled in the art, for purposes of obtaining antisera containing polyclonal antibodies or hybridoma lines secreting monoclonal antibodies.

Monoclonal antibodies can be prepared by standard techniques, given the teachings contained herein. Such techniques are disclosed, for example, in U.S. Patent No. 4,271,145 and U.S. Patent No. 4,196,265. Briefly, an animal is immunized with the immunogen. Hybridomas are prepared by fusing spleen cells from the immunized animal with myeloma cells. The fusion products are screened for those producing antibodies that 25 bind to the immunogen. The positive hybridomas clones are isolated, and the monoclonal antibodies are recovered from those clones.

Immunization regimens for production of both polyclonal and monoclonal antibodies are well known in the art. The immunogen may be injected by any of a number of routes, including subcutaneous, intravenous, intraperitoneal, intradermal, intramuscular, 30 mucosal, or a combination of these. The immunogen may be injected in soluble form, aggregate form, attached to a physical carrier, as a gel slice, or mixed with an adjuvant, using methods and materials well known in the art. The antisera and antibodies may be purified using column chromatography methods well known to those of skill in the art.

The antibodies may also be used as probes for identifying clones in 35 expression libraries that have or may have inserts encoding one or more PMP polypeptides

or fragments thereof. The antibodies, PMP polypeptides or PMP-derived polypeptides may also be used in immunoassays (e.g., ELISA, RIA, Western Blots) to specifically detect and/or quantitate *Chlamydia* or anti-*Chlamydia* antibody in biological specimens. Anti-PMP antibodies of the invention specifically bind PMP polypeptide from *Chlamydia* trachomatis, *Chlamydia pneumoniae*, *Chlamydia pecorum*, and/or *Chlamydia psittaci*. Thus anti-PMP antibodies can be used to diagnose *Chlamydia* infections.

The antibodies of the invention, including but not limited to those that are cytotoxic, cytostatic, or neutralizing, may also be used in passive immunization to prevent or attenuate *Chlamydia* infections of animals, including humans. As used herein, a cytotoxic antibody is one that enhances opsonization and/or complement killing of the bacterium bound by the antibody. As used herein, neutralizing antibody is one that reduces the infectivity of the *Chlamydia* and/or blocks binding of *Chlamydia* to a target cell. An effective concentration of polyclonal or monoclonal antibodies raised against the immunogens of the invention may be administered to a host to achieve such effects. The exact concentration of the antibodies administered will vary according to each specific antibody preparation, but may be determined using standard techniques well known to those of ordinary skill in the art. Administration of the antibodies may be accomplished using a variety of techniques, including, but not limited to those described in Section 5.7 for the delivery of vaccines.

Another aspect of the invention is directed to antisera raised against an antigenic or immunogenic composition of the invention, and antibodies present in the antisera that specifically bind a PMP protein or a fragment or analogue thereof.

Preferably, the antibodies bind a PMP polypeptide having the amino acid sequence selected from the group consisting of SEQ ID NOs.:2 and 4-34 or a PMP-derived polypeptide. Also included are monoclonal antibodies that specifically bind a PMP polypeptide having the amino acid sequence selected from the group consisting of SEQ ID NOs.:2 and 4-34 or a PMP-derived polypeptide. The term "antibodies" is intended to include all forms, such as but not limited to polyclonal, monoclonal, purified IgG, IgM, or IgA antibodies and fragments thereof, including but not limited to antigen binding fragments such as Fv, single chain Fv (scFv), F(ab.)₂, Fab, and F(ab)' fragments (Harlow and Lane, 1988, Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory Press); single chain antibodies (U.S. Patent No. 4,946,778) and complementary determining regions (CDR), (see Verhoeyen and Winter, in Molecular Immunology 2ed., by B.D. Hames and D.M. Glover, IRL Press, Oxford University Press, 1996, at pp. 283-325), etc.

A further aspect of the invention are chimeric or humanized antibodies (Morrison et al., 1984, Proc. Nat'l Acad. Sci. USA 81:6851; Reichmann et al. Nature 332:323: U.S. Patent Nos. 5,225,539; 5,585,089; and 5,530,101; Neuberger et al., 1984, Nature 81:6851 Riechmann et al., 1988, Nature 332:323; U.S. Patent Nos. 5,225,539; 5,585,089; and 5,530,101) in which one or more of the antigen binding regions of the anti-PMP antibody is introduced into the framework region of a heterologous (e.g., human) antibody. The chimeric or humanized antibodies of the invention are less antigenic in humans than non-human antibodies but have the desired antigen binding and other

activities, including but not limited to neutralizing activity, cytotoxic activity, opsonizing

10 activity or protective activity.

In a preferred embodiment, the antibodies of the invention are human antibodies. Human antibodies may be isolated, for example, from human immunoglobulin libraries (see, e.g., PCT publications WO 98/46645, WO 98/50433, WO 98/24893, WO 98/16054, WO 96/34096, WO 96/33735, and WO 91/10741) by, preferably, phage display techniques (see, e.g., Brinkman et al., J. Immunol. Methods 182:41-50 (1995); Ames et al., J. Immunol. Methods 184:177-186 (1995); Kettleborough et al., Eur. J. Immunol. 24:952-958 (1994); Persic et al., Gene 187 9-18 (1997); Burton et al., Advances in Immunology 57:191-280 (1994); PCT application No. PCT/GB91/01134; PCT publications WO 90/02809; WO 91/10737; WO 92/01047; WO 92/18619; WO 93/11236; WO 95/15982;

- WO 95/20401; and U.S. Patent Nos. 5,698,426; 5,223,409; 5,403,484; 5,580,717; 5,427,908; 5,750,753; 5,821,047; 5,571,698; 5,427,908; 5,516,637; 5,780,225; 5,658,727; 5,733,743 and 5,969,108; each of which is incorporated herein by reference in its entirety. Human antibodies may also be generated from animals transgenic for one or more human immunoglobulin and that do not express endogenous immunoglobulins, see, *e.g.*, U.S.
- 25 Patent No. 5,939,598 by Kucherlapati et al.

A further aspect of the invention is T-cells specific for *Chlamydia*, Chlamydial EB, RBs, *Chlamydia* infected cells or antigen presenting cells displaying Chlamydial antigens. T-cell preparations enriched for T-cells specific for PMP or PMP-derived polypeptides can be produced or isolated by methods known in the art (See section 30 5.8).

5.5 NUCLEIC ACIDS ENCODING THE PMP POLYPEPTIDE AND PMP DERIVED POLYPEPTIDES

The isolated nucleic acids of the present invention, including DNA and RNA, comprising a sequence encoding the PMP protein or PMP-derived polypeptide

thereof, may be synthesized using methods known in the art, such as using conventional chemical approaches or polymerase chain reaction (PCR) amplification using convenient pairs of oligonucleotide primers and ligase chain reaction using a battery of contiguous oligonucleotides. The sequences also allow for the identification and cloning of the PMP protein gene from any species or serovar of *Chlamydia*, for instance for screening Chlamydial genomic libraries or expression libraries as described below.

In a particular embodiment, the PMP polypeptide comprises an amino acid sequence of either SEQ ID NO.:2 or 4 and the nucleic acids comprise nucleotide sequences encoding said amino acid sequences. Particularly preferred fragments of PMP have 8 or more amino acids of the amino acid sequences of SEQ ID NOs.:2 or 4, or sequences substantially homologous thereto, and the invention encompasses nucleic acids comprising nucleotide sequences encoding said amino acid sequences. In another particular embodiment, the PMP polypeptide is encoded by the nucleotide sequence of SEQ ID NOs.:1 or 3, with particularly preferred fragments having a nucleotide sequence of SEQ ID NOs.:36-65, or sequences substantially homologous thereto.

The term "isolated nucleic acid" or "isolated nucleic acid molecule" is defined as a nucleic acid molecule or removed from the environment in which it naturally occurs. For example, a naturally-occurring DNA molecule present in the genome of a living bacteria or as part of gene bank is not isolated, but the same molecule separated from the remaining part of the bacterial genome, as a result of, *e.g.*, a cloning event (amplification) is isolated. Typically, an isolated DNA molecule is free from DNA regions (*e.g.*, coding regions) with which it is immediately contiguous at the 5' or 3' end, in the naturally occurring genome. Such isolated nucleic acids or nucleic acid molecules could be part of a vector or a composition and still be isolated in that such a vector or composition is not part of its natural environment. However, "isolated nucleic acid" or "isolated nucleic acid molecule" does not include a nucleic acid that is part of a recombinant library.

Nucleic acids of the present invention can be single or double stranded. The invention also provides nucleic acids hybridizable to or complementary to SEQ ID NO.:1 or 3 or fragments thereof, as well as polypeptides encoded by these nucleic acids. In specific aspects, nucleic acids are provided which comprise a sequence fully complementary to or complementary to at least 10, 15, 25, 50, 100, 200, 250, 300, 350, 400, 450, 500, 550, 600, 650, 700, 750, 800, 850, 900, 950, 1000, 1100, 1200, 1300, 1400 or 1500 contiguous nucleotides of a nucleic acid encoding a PMP polypeptide or an PMP-derived polypeptide. In a specific embodiment, a nucleic acid which is hybridizable to a nucleic acid encoding a PMP polypeptide (e.g., having a nucleotide sequence of SEQ ID NO.:1 or 3), or to a nucleic

acid encoding a PMP-derived polypeptide, under conditions of low, moderate or high stringency is provided.

By way of example and not limitation, procedures using such conditions of low stringency are as follows (see also Shilo and Weinberg, 1981, Proc. Natl. Acad. Sci.

5 USA 78:6789-6792): Filters containing DNA are pretreated for 6 h at 40°C in a solution containing 35% formamide, 5X SSC, 50 mM Tris-HCl (pH 7.5), 5 mM EDTA, 0.1% PVP, 0.1% Ficoll, 1% BSA, and 500 μg/ml denatured salmon sperm DNA. Hybridizations are carried out in the same solution with the following modifications: 0.02% PVP, 0.02% Ficoll, 0.2% BSA, 100 μg/ml salmon sperm DNA, 10% (wt/vol) dextran sulfate, and 5-20 X 10 cpm ³²P-labeled probe is used. Filters are incubated in hybridization mixture for 18-20 h at 40°C, and then washed for 1.5 h at 55°C in a solution containing 2X SSC, 25 mM Tris-HCl (pH 7.4), 5 mM EDTA, and 0.1% SDS. The wash solution is replaced with fresh solution and incubated an additional 1.5 h at 60°C. Filters are blotted dry and exposed for autoradiography. If necessary, filters are washed for a third time at 65-68°C and re-exposed to film. Other conditions of low stringency which may be used are well known in the art (e.g., as employed for cross-species hybridizations).

In another specific embodiment, a nucleic acid which is hybridizable to a nucleic acid encoding a PMP polypeptide or a PMP-derived polypeptide under conditions of high stringency is provided. By way of example and not limitation, procedures using such conditions of high stringency are as follows: Prehybridization of filters containing DNA is carried out for 8 h to overnight at 65°C in buffer composed of 6X SSC, 50 mM Tris-HCl (pH 7.5), 1 mM EDTA, 0.02% PVP, 0.02% Ficoll, 0.02% BSA, and 500 μg/ml denatured salmon sperm DNA. Filters are hybridized for 48 h at 65°C in prehybridization mixture containing 100 μg/ml denatured salmon sperm DNA and 5-20 X 106 cpm of ³²P-labeled probe. Washing of filters is done at 37°C for 1 h in a solution containing 2X SSC, 0.01% PVP, 0.01% Ficoll, and 0.01% BSA. This is followed by a wash in 0.1X SSC at 50°C for 45 min before autoradiography. Other conditions of high stringency which may be used are well known in the art.

In another specific embodiment, a nucleic acid which is hybridizable to a nucleic acid encoding a PMP polypeptide or a PMP-derived polypeptide under conditions of moderate stringency is provided.

Various other stringency conditions which promote nucleic acid hybridization can be used. For example, hybridization in 6X SSC at about 45°C, followed by washing in 2X SSC at 50°C may be used. Alternatively, the salt concentration in the wash step can range from low stringency of about 5X SSC at 50°C, to moderate stringency

of about 2X SSC at 50°C, to high stringency of about 0.2X SSC at 50°C. In addition, the temperature of the wash step can be increased from low stringency conditions at room temperature, to moderately stringent conditions at about 42°C, to high stringency conditions at about 65°C. Other conditions include, but are not limited to, hybridizing at 68°C in 0.5M NaHPO₄ (pH7.2)/1 mM EDTA/7% SDS, or hybridization in 50% formamide/0.25M NaHPO₄ (pH 7.2)/.25 M NaCl/1 mM EDTA/7% SDS; followed by washing in 40 mM NaHPO₄ (pH 7.2)/1 mM EDTA/5% SDS at 42°C or in 40 mM NaHPO₄ (pH7.2) 1 mM EDTA/1% SDS at 50°C. Both temperature and salt may be varied, or alternatively, one or the other variable may remain constant while the other is changed.

Low, moderate and high stringency conditions are well known to those of skill in the art, and will vary predictably depending on the base composition of the particular nucleic acid sequence and on the specific organism from which the nucleic acid sequence is derived. For guidance regarding such conditions see, for example, Sambrook et al., 1989, Molecular Cloning, A Laboratory Manual, Second Edition, Cold Spring Harbor Press, N.Y., pp. 9.47-9.57; and Ausubel et al., 1989, Current Protocols in Molecular Biology, Green Publishing Associates and Wiley Interscience, N.Y.

In the preparation of genomic libraries, DNA fragments are generated, some of which will encode parts or the whole of a *Chlamydia* PMPE or PMPI protein. The DNA may be cleaved at specific sites using various restriction enzymes. Alternatively, one may use DNase in the presence of manganese to fragment the DNA, or the DNA can be physically sheared, as for example, by sonication. The DNA fragments can then be separated according to size by standard techniques, including but not limited to, agarose and polyacrylamide gel electrophoresis, column chromatography and sucrose gradient centrifugation. The DNA fragments can then be inserted into suitable vectors, including but not limited to plasmids, cosmids, bacteriophages lambda or T₄, bacmids and yeast artificial chromosome (YAC). (See, for example, Sambrook et al., 1989, Molecular Cloning, A Laboratory Manual, 2d Ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York; Glover, D.M. (ed.), 1985, DNA Cloning: A Practical Approach, MRL Press, Ltd., Oxford, U.K. Vol. I, II.) The genomic library may be screened by nucleic acid hybridization to labeled probe (Benton and Davis, 1977, Science 196:180; Grunstein and Hogness, 1975, Proc. Natl. Acad. Sci. U.S.A.. 72:3961).

The genomic libraries may be screened with labeled degenerate oligonucleotide probes corresponding to the amino acid sequence of any peptide of PMP protein using optimal approaches well known in the art. Any probe used preferably is 15 nucleotides or longer.

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The term "probe" as used herein refers to DNA (preferably single stranded) or RNA molecules that hybridize under stringent conditions as defined above, to nucleic acids having sequences homologous to SEQ ID NO.:1 or SEQ ID NO.:3 or to a complementary or anti-sense sequence thereof. Generally, probes are significantly shorter than full-length sequences shown in SEQ ID NO.:1 or 3.

For example, they can contain from about 5 to about 100 nucleotides preferably from about 10 to about 80 nucleotides. In particular, probes have sequences that are at least 75% preferably at lest 85%, and more preferably 95%, homologous to a portion of a sequence of SEQ ID NO.:1 or 3, or complementary to such sequences. Probes can contain modified bases such as inosine, methyl-5-deoxycytidine, deoxyuridine, dimethylamino-5-deoxyuridine, or diamino-2,6 purine.

Clones in libraries with insert DNA encoding a PMP protein or a PMP-derived polypeptides will hybridize to one or more of the degenerate oligonucleotide probes. Hybridization of such oligonucleotide probes to genomic libraries is carried out using methods known in the art. For example, hybridization with the two above-mentioned oligonucleotide probes may be carried out in 2X SSC, 1.0% SDS at 50°C and washed using the same conditions.

In yet another aspect, clones of nucleotide sequences encoding a part or the entire PMP protein or PMP-derived polypeptide may also be obtained by screening

20 *Chlamydia* expression libraries. For example, *Chlamydia* DNA or *Chlamydia* cDNA generated from RNA is isolated and random fragments are prepared and ligated into an expression vector (*e.g.*, a bacteriophage, plasmid, phagemid or cosmid) such that the inserted sequence in the vector is capable of being expressed by the host cell into which the vector is then introduced. Various screening assays can then be used to select for the

25 expressed PMP-protein or PMP-derived polypeptides. In one embodiment, the various anti-PMP antibodies of the invention can be used to identify the desired clones using methods known in the art. See, for example, Harlow and Lane, 1988, <u>Antibodies: A Laboratory Manual</u>, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, Appendix IV. Clones or plaques from the library are brought into contact with the antibodies to identify those clones that bind.

In an embodiment, colonies or plaques containing DNA that encodes a PMP protein or PMP-derived polypeptide could be detected using DYNA Beads according to Olsvick et al., 29th ICAAC, Houston, Tex. 1989, incorporated herein by reference. Anti-PMP antibodies are crosslinked to DYNA Beads M280, and these antibody-containing beads are used to adsorb to colonies or plaques expressing a PMP protein or a PMP derived

polypeptide. Colonies or plaques expressing a PMP protein or a PMP derived polypeptide are identified as any of those that bind the beads.

Alternatively, the anti-PMP antibodies can be nonspecifically immobilized to a suitable support, such as silica or CeliteTM resin. This material is used to adsorb to bacterial colonies expressing a PMP protein or a PMP derived polypeptide as described in the preceding paragraph.

In another aspect, PCR amplification may be used to produce substantially pure DNA encoding a part of or the whole of a PMP protein from *Chlamydia* genomic DNA. Oligonucleotide primers, degenerate or otherwise, corresponding to known PMP protein sequences can be used as primers.

In particular embodiments, an oligonucleotide encoding a portion of SEQ ID NO.:2 or 4 may be used as the 5' primer. For fragment examples, a 5' primer may be made from any one of the nucleotide sequences of SEQ ID NO.:66 or 69 or any portion thereof. For 3' primers, a nucleotide sequence of SEQ ID NO.:67 or 70 or any portion thereof may be used.

As examples, an oligonucleotide encoding the N-terminal primer, and together with a 3' reverse PCR oligonucleotide complementary to an internal, downstream protein coding sequence may be used to amplify an N-terminal-specific PMP DNA fragment. Alternatively, an oligonucleotide encoding an internal PMP coding sequence may be used as the 5' forward PCR primer together with a 3' reverse PCR oligonucleotide complementary to downstream, internal PMP protein coding sequences may be used to PCR amplify an internal PMP specific DNA fragment. Alternatively, the forward primer can be combined together with an oligonucleotide complementary to the C-terminal PMP coding region to PCR amplify the PMP ORF. These PMP specific PCR products can be cloned into appropriate expression vectors to direct the synthesis of all or part of the PMP polypeptide as distinct proteins or fusion proteins. Alternatively, these PMP specific PCR products can be appropriately labeled and used as hybridization probes to identify all or part of the PMP gene from genomic DNA libraries.

PCR can be carried out, *e.g.*, by use of a Perkin-Elmer thermal cycler and 30 Taq polymerase (Gene Amp[™]). One can choose to synthesize several different degenerate primers, for use in the PCR reactions. It is also possible to vary the stringency of hybridization conditions used in priming the PCR reactions, to allow for greater or lesser degrees of nucleotide sequence similarity between the degenerate primers and the corresponding sequences in *Chlamydia* DNA. After successful amplification of a segment of the sequence encoding a PMP protein, that segment may be molecularly cloned and

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sequenced, and utilized as a probe to isolate a complete genomic clone. This, in turn, will permit the determination of the gene's complete nucleotide sequence, the analysis of its expression, and the production of its protein product for functional analysis, as described infra.

Once a PMP polypeptide coding sequence has been isolated from one Chlamydia species, strain, or cultivar, it is possible to use the same approach to isolate PMP polypeptide coding sequences from other Chlamydia species, strains and cultivars. It will be recognized by those skilled in the art that the DNA or RNA sequence encoding PMP polypeptide (or fragments thereof) of the invention can be used to obtain other DNA or 10 RNA sequences that hybridize with it under conditions of moderate to high stringency, using general techniques known in the art (see supra). Hybridization with PMP sequence from one Chlamydia strain or cultivar under high stringency conditions will identify the corresponding sequence from other strains and cultivars. High stringency conditions vary with probe length and base composition. The formulae for determining such conditions are 15 well known in the art. See Sambrook et al., 1989, Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Press, NY, Chapter 11. As an example, high stringency hybridization conditions as applied to probes of greater than 300 bases in length involve a final wash in 0.1X SSC/0.1% SDS at 68°C for at least 1 hour (Ausubel, et al., Eds., 1989, Current Protocols in Molecular Biology, Vol. I, Greene Publishing Associates, Inc and John 20 Wiley & Sons, Inc. New York, at page 2.10.2). See also, the description of high stringency conditions, supra.

One skilled in the art would be able to identify complete clones of a PMP polypeptide coding sequence using approaches well known in the art. The extent of the PMP polypeptide coding sequence contained in an isolated clone may be ascertained by 25 sequencing the cloned insert and comparing the deduced size of the polypeptide encoded by the open reading frames (ORFs) with that of the PMP polypeptide and/or by comparing the deduced amino acid sequence with that of known amino acid sequence of the purified PMP polypeptide. Where a partial clone of the PMP polypeptide coding sequence has been isolated, complete clones may be isolated by using the insert of the partial clone as a 30 hybridization probe. Alternatively, a complete PMP polypeptide coding sequence can be reconstructed from overlapping partial clones by splicing their cloned PMP inserts together.

Complete clones may be any that an ORF with a deduced amino acid sequence matching or substantially homologous to that of the PMP polypeptide or, where the complete amino acid sequence of the latter is not available, matching or substantially 35 homologous to that of a peptide fragment of a PMP polypeptide and having a molecular

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weight corresponding to that of the PMP polypeptide. Further, complete clones may be identified by the ability of their inserts, when placed in an expression vector, to produce a polypeptide that binds antibodies specific to the amino-terminus of the PMP polypeptide and antibodies specific to the carboxyl-terminus of the PMP polypeptide.

Nucleic acids encoding PMP-derived polypeptides and PMP fusion proteins may be produced by methods well known in the art. In one aspect, nucleic acids encoding PMP-derived polypeptides can be derived from PMP polypeptide coding sequences by recombinant DNA methods in view of the teachings disclosed herein. For example, the coding sequence of a PMP polypeptide may be altered creating amino acid substitutions that 10 will not affect the immunogenicity of the PMP polypeptide or which may improve its immunogenicity, such as conservative or semi-conservative substitutions as described above. Various methods may be used, including but not limited to, oligonucleotide directed, site specific mutagenesis. This and other techniques known in the art may be used to create single or multiple mutations, such as replacements, insertions, deletions, and 15 transpositions, for example, as described in Botstein and Shortle, 1985, Science 229:1193-1210.

Further, nucleic acids containing PMP polypeptide coding sequences may be truncated by restriction enzyme or exonuclease digestions. Heterologous coding sequences may be added to the PMP polypeptide coding sequence by ligation or PCR amplification. 20 Moreover, DNA encoding the whole or a part of PMP-derived polypeptide may be

synthesized chemically or using PCR amplification based on the known or deduced amino acid sequence of the PMP polypeptide and any desired alterations to that sequence.

The identified and isolated DNA containing the PMP polypeptide or PMPderived polypeptide coding sequence can be inserted into an appropriate cloning vector. A 25 large number of vector-host systems known in the art may be used. The term "host" as used herein and in the claims refers to either in vivo in an animal or in vitro in mammalian cell cultures.

Possible vectors include, but are not limited to, plasmids and modified viruses, but the vector system must be compatible with the host cell used. Such vectors 30 include, but are not limited to, bacteriophage such as lambda derivatives, or plasmids such as pET, pBAD, pTrcHis, pBR322 or pUC plasmid derivatives. The insertion into a cloning vector can, for example, be accomplished by ligating the DNA fragment into a cloning vector which has complementary cohesive termini. However, if the complementary restriction sites used to fragment the DNA are not present in the cloning vector, the ends of 35 the DNA molecules may be enzymatically modified. Alternatively, any site desired may be produced by ligating nucleotide sequences (linkers) onto the DNA termini; these ligated linkers may comprise specific chemically synthesized oligonucleotides encoding restriction endonuclease recognition sequences. In an alternative method, the cleaved DNA may be modified by homopolymeric tailing. Recombinant molecules can be introduced into host cells via transformation, transfection, infection, electroporation, etc., so that many copies of the gene sequence are generated.

In an alternative method, the desired DNA containing a PMP polypeptide or PMP-derived polypeptide coding sequence may be identified and isolated after insertion into a suitable cloning vector in a "shot gun" approach. Enrichment for the desired sequence, for example, by size fractionation, can be done before insertion into the cloning vector.

In specific embodiments, transformation of host cells with recombinant DNA molecules that contain a PMP polypeptide or PMP-derived polypeptide coding sequence enables generation of multiple copies of such coding sequence. Thus, the coding sequence may be obtained in large quantities by growing transformants, isolating the recombinant DNA molecules from the transformants and, when necessary, retrieving the inserted coding sequence from the isolated recombinant DNA.

The nucleotide sequences encoding the PMP polypeptides of the present invention are useful for their ability to selectively form duplex molecules with

20 complementary stretches of other protein genes. Depending on the application, a variety of hybridization conditions may be employed to achieve hybridization with varying sequence identities. In specific aspects, nucleic acids are provided which comprise a sequence complementary to at least 10, 15, 25, 50, 100, 200 or 250 nucleotides of the PMP protein coding nucleic acid molecule. In specific embodiments, nucleic acids which hybridize to a PMP protein nucleic acid (*e.g.*, having a nucleotide sequence of SEQ ID NO.:1 or 3) under annealing conditions are provided.

For a high degree of selectivity, relatively stringent conditions are used to form the duplexes, such as, by way of example and not limitation, low salt and/or high temperature conditions, such as provided by 0.02 M to 0.15 M NaCl at temperatures of between about 50°C to 70°C. For some applications, less stringent hybridization conditions are required, by way of example and not limitation, such as 0.15 M to 0.9 M salt, at temperatures ranging from between about 20°C to 55°C. Hybridization conditions can also be rendered more stringent by the addition of increasing amounts of formamide, to destabilize the hybrid duplex. Thus, particular hybridization conditions can be readily manipulated, and will generally be a method of choice depending on the desired results.

5.6 <u>RECOMBINANT PRODUCTION OF PMP POLYPEPTIDE AND PMP-DERIVED POLYPEPTIDES</u>

In accordance with this invention, it is preferred to make the *Chlamydia*5 protein of the present invention by recombinant methods, particularly when the naturally occurring protein as isolated from a culture of a species of *Chlamydia* may include trace amounts of toxic materials or other contaminants. This problem can be avoided by using protein recombinantly produced in heterologous systems which can be isolated from the host in a manner to minimize contaminants in the isolated material. In this case, the PMP proteins are produced by an appropriate host cell that has been transformed by a DNA molecule that codes for the polypeptide.

The nucleic acids encoding the PMP polypeptides or PMP-derived polypeptides of the invention can be inserted into an appropriate expression vector, *i.e.*, a vector that contains the necessary elements for the transcription and translation of the inserted polypeptide-coding sequence. The nucleotide sequences encoding the PMP polypeptides or PMP-derived polypeptides are inserted into the vectors in a manner such that they will be expressed under appropriate conditions (*e.g.*, in proper orientation and correct reading frame). The recombinant expression vector also comprises an "expression means". The term "expression means" refers to elements of a vector which are necessary for transcription and translation of the nucleic acid encoding the protein, including but not limited to promoter/enhancer elements, a replication site, an RNA polymerase binding sequence, a ribosomal binding sequence, sequences which are capable of providing phenotype selection (*e.g.*, ampicillin or tetracycline resistance) and replicon and control sequences that can be used to transform host cells. The expression means is operatively coupled to the nucleic acid molecule encoding the PMP protein by linking the inserted nucleic acid molecule into the expression vector.

Promoter/enhancer elements which may be used to control expression of inserted sequences include, but are not limited to the SV40 early promoter region (Bernoist and Chambon, 1981, Nature 290:304-310), the promoter contained in the 3' long terminal repeat of Rous sarcoma virus (Yamamoto et al., 1980, Cell 22:787-797), the herpes thymidine kinase promoter (Wagner et al., 1981, Proc. Natl. Acad. Sci. U.S.A.. 78:1441-1445), the regulatory sequences of the metallothionein gene (Brinster et al., 1982, Nature 296:39-42) for expression in animal cells; the promoters of lactamase (Villa-Kamaroff et al., 1978, Proc. Natl. Acad. Sci. U.S.A.. 75:3727-3731), tac (DeBoer et al., 1983, Proc.

35 Natl. Acad. Sci. U.S.A. 80:21-25), or trc for expression in bacterial cells (see also "Useful

proteins from recombinant bacteria" in <u>Scientific American</u>, 1980, 242:74-94); the nopaline synthetase promoter region or the cauliflower mosaic virus 35S RNA promoter (Gardner et al., 1981, <u>Nucl. Acids Res.</u> 9:2871), and the promoter of the photosynthetic enzyme ribulose biphosphate carboxylase (Herrera-Estrella et al., 1984, <u>Nature</u> 310:115-120) for expression in plant cells; Gal4 promoter, the ADC (alcohol dehydrogenase) promoter, PGK (phosphoglycerol kinase) promoter, alkaline phosphatase promoter for expression in yeast or other fungi.

Depending on the host-vector system utilized, any one of a number of suitable transcription and translation elements may be used. In a preferred embodiment, a chimeric protein comprising a PMP protein or PMP-derived polypeptide sequence and a pre and/or pro sequence of the host cell is expressed. In other preferred embodiments, a chimeric protein comprising a PMP protein or PMP derived polypeptide sequence fused with, for example, an affinity purification peptide, including but not limited to, maltose binding protein, glutathione-S-transferase, thioredoxin or histidine tag, is expressed. In further preferred embodiments, a chimeric protein comprising a PMP protein or PMP-derived polypeptide sequence and a useful immunogenic peptide or protein is expressed.

Any method known in the art for inserting DNA fragments into a vector may be used to construct expression vectors containing a PMP or PMP-derived polypeptide encoding nucleic acid molecule consisting of appropriate transcriptional/translational control signals and the polypeptide coding sequences. These methods may include *in vitro* recombinant DNA and synthetic techniques and *in vivo* recombinantation (genetic recombination).

Methods of introducing exogenous DNA into yeast hosts include either the transformation of spheroplasts or of intact yeast cells treated with alkali cations.

- Transformation procedures usually vary with the yeast species to be transformed. See *e.g.*, Kurtz et al. (1986) Mol. Cell. Biol. 6:142; Kunze et al. (1985) J. Basic Microbiol. 25:141; for Candida, Gleeson et al. (1986) J. Gen. Microbiol. 132:3459; Roggenkamp et al. (1986) Mol. Gen. Genet. 202:302; for Hansenula; Das et al. (1984) J. Bacteriol. 158:1165; De Louvencourt et al. (1983) J. Bacteriol. 154:1165; Van den Berg et al. (1990)
- 30 <u>Bio/Technology</u> 8:135; for *Kluyveromyces*; Cregg et al. (1985) <u>Mol. Cell. Biol.</u> 5:3376; Kunze et al. (1985) <u>J. Basic Microbiol.</u> 25:141; U.S. Patent No. 4,837,148 and U.S. Patent No. 4,929,555; for *Pichia*; Hinnen et al. (1978) <u>Proc. Natl. Acad. Sci.</u> USA 75;1929; Ito et al. (1983) <u>J. Bacteriol</u>. 153:163; for *Saccharomyces*; Beach et al. (1981) <u>Nature</u> 300:706; for *Schizosaccharomyces*; Davidow et al. (1985) <u>Curr. Genet.</u> 10:39.

Expression vectors containing PMP polypeptide or PMP-derived polypeptide coding sequences can be identified by three general approaches: (a) nucleic acid hybridization; (b) presence or absence of "marker" gene functions; and (c) expression of inserted sequences. In the first approach, the presence of a foreign gene inserted into an expression vector can be detected by nucleic acid hybridization using probes comprising sequences that are homologous to the inserted PMP polypeptide or PMP-derived polypeptide coding sequence. In the second approach, the recombinant vector/host system can be identified and selected based upon the presence or absence of certain "marker" gene functions (*e.g.*, thymidine kinase activity, resistance to antibiotics, transformation phenotype, occlusion body formation in baculovirus, etc.) caused by the insertion of foreign genes into the vector.

For example, *E. coli* may be transformed using pBR322 which contains genes for ampicillin and tetracycline resistance. If the PMP polypeptide or PMP-derived polypeptide coding sequence is inserted within the marker gene sequence of the vector, recombinants containing the insert can be identified by the absence of the marker gene function. In the third approach, recombinant expression vectors can be identified by assaying the foreign gene product expressed by the recombinant. Such assays can be based, for example, on the physical or functional properties of PMP polypeptide or PMP-derived polypeptide in *vitro* assay systems, *e.g.*, binding of a His tag to a column, binding to a PMP ligand or receptor or binding with anti-PMP antibodies of the invention.

Commercially available vectors for expressing heterologous proteins in bacterial hosts include but are not limited to pZERO, pTrc99A, pUC19, pUC18, pKK223-3, pEX1, pCAL, pET, pSPUTK, pTrxFus, pFastBac, pThioHis, pTrcHis, pTrcHis2, and pLEx. For example, the phage in lambda GEMTM-11 may be utilized in making recombinant phage vectors which can be used to transform host cells, such as *E. coli* LE392. In a preferred embodiment, the vector is pQE30 or pBAD/ThioE, which can be used transform host cells, such as *E. coli*.

Expression and transformation vectors for transformation into many yeast strains are available. For example, expression vectors have been developed for, the following yeasts: *Candida albicans*, Kurtz, et al. (1986) Mol. Cell. Biol. 6:142; *Candida maltosa*, Kunze, et al. (1985) J. Basic Microbiol. 25:141; *Hansenula polymorpha*, Gleeson, et al. (1986) J. Gen. Microbiol. 132:3459; Roggenkamp et al. (1986) Mol. Gen. Genet. 202:302; *Kluyveromyces fragilis*, Das, et al. (1984) J. Bacteriol. 158:1165; *Kluyveromyces lactis*, De Louvencourt et al. (1983) J. Bacteriol. 154:737; Van den Berg, et al. (1990) Bio/Technology 8:135; *Pichia quillerimondii*, Kunze et al. (1985) J. Basic Microbiol.

25:141; *Pichia pastoris*, Cregg, et al. (1985) Mol. Cell. Biol. 5:3376, U.S. Patent No. 4,837,148 and U.S. Patent No. 4,929,555; *Saccharomyces cerevisiae*, Hinnen et al. (1978) Proc. Natl. Acad. Sci. USA 75:1929, Ito et al. (1983) J. Bacteriol. 153:163; *Schizosaccharomyces pombe*, Beach et al. (1981) Nature 300:706; and *Yarrowia lipolytica*, Davidow, et al. (1985) Curr. Genet. 10:380-471, Gaillardin, et al. (1985) Curr. Genet. 10:49.

A variety of host-vector systems may be utilized to express the polypeptide-coding sequence. These include but are not limited to mammalian cell systems infected with virus (e.g., vaccinia virus, adenovirus, etc.); insect cell systems infected with virus 10 (e.g., baculovirus); microorganisms such as yeast containing yeast vectors, or bacteria transformed with bacteriophage DNA, plasmid DNA, or cosmid DNA, plant cells or transgenic plants.

Hosts that are appropriate for expression of nucleic acid molecules of the present invention, fragments, analogues or variants thereof, may include *E. coli*, *Bacillus* species, *Haemophilus*, fungi, yeast, such as *Saccharomyces*, *Pichia*, *Bordetella*, or *Candida*, or the baculovirus expression system may be used. Preferably, the host cell is a yeast or bacterium.

Particularly desirable hosts for expression in this regard include Gram positive bacteria which do not have LPS and are, therefore endotoxin free. Most preferably 20 the bacterium is *E. coli*, *B. subtilis* or *Salmonella*.

In addition, a host cell strain may be chosen which modulates the expression of the inserted sequences, or modifies and processes the gene product in the specific fashion desired. Expression from certain promoters can be elevated in the presence of certain inducers; thus, expression of the genetically engineered PMP polypeptide or PMP-derived polypeptide may be controlled. Furthermore, different host cells have characteristic and specific mechanisms for the translational and post-translational processing and modification of proteins. Appropriate cell lines or host systems can be chosen to ensure the desired modification and processing of the foreign protein expressed.

Once a suitable host system and growth conditions are established,
recombinant expression vectors can be propagated and prepared in quantity. Upon
expression, a recombinant polypeptide of the invention is produced and can be recovered in
a substantially purified from the cell paste, the cell extract or from the supernatant after
centrifugation of the recombinant cell culture using techniques well known in the art.

For instance, the recombinant polypeptide can be purified by antibody-based affinity purification, preparative gel electrophoresis, or affinity purification using tags (*e.g.*, 6X histidine tag) included in the recombinant polypeptide. (See, Section 5.3 *supra*).

5 5.7 <u>COMPOSITIONS</u>

The present invention also provides therapeutic and prophylactic compositions, which may be antigenic compositions, and preferably immunogenic compositions, including vaccines, for use in the treatment or prevention of Chlamydia infections of animals, including mammals and birds, and more specifically rodents and 10 primates, including humans. Preferred immunogenic compositions include vaccines for use in humans. The antigenic, preferably immunogenic, compositions of the present invention can be prepared by techniques known to those skilled in the art and comprise, for example, an immunologically effective amount of any of the PMP immunogens disclosed in Sections 5.1. or 5.2, optionally in combination with or fused to or conjugated to one or more other 15 immunogens, including lipids, phospholipids, carbohydrates, lipopolysaccharides, inactivated or attenuated whole organisms and other proteins, of Chlamydia origin or other bacterial origin, a pharmaceutically acceptable carrier, optionally an appropriate adjuvant, and optionally other materials traditionally found in vaccines. In one embodiment, the invention provides a cocktail vaccine comprising several immunogens, which has the 20 advantage that immunity against one or several strains of a single pathogen or one or several pathogens can be obtained by a single administration. Examples of other immunogens include, but are not limited to, those used in the known DPT vaccines, HMWP of C. trachomatis, MOMP of C. trachomatis, entire organisms or subunits therefrom of Chlamydia, Neisseria, HIV, Haemophilus influenzae, Moraxella catarrhalis, Human 25 papilloma virus, Herpes simplex virus, Haemophilus ducreyi, Treponema pallidium, Candida albicans and Streptococcus pneumoniae, etc.

The term "immunogenic amount" is used herein to mean an amount sufficient to induce an immune response to produce antibodies, T-cells, and/or cytokines and other cellular immune response components. Preferably, the immunogenic composition is one that elicits an immune response sufficient to prevent *Chlamydia* infections or to attenuate the severity of any preexisting or subsequent *Chlamydia* infection. An immunogenic amount of the immunogen to be used in the vaccine is determined by means known in the art in view of the teachings herein. The exact concentration will depend upon the specific immunogen to be administered, but can be determined by using standard

techniques well known to those skilled in the art for assaying the development of an immune response.

The vaccine compositions of the invention elicit an immune response in a subject. Compositions which induce antibodies, including anti-PMP protein antibodies and antibodies that are opsonizing or bactericidal are one aspect of the invention. In preferred non-limiting, embodiments of the invention, an effective amount of a composition of the invention produces an elevation of antibody titer to at least three times the antibody titer prior to administration. In a preferred, specific, non-limiting embodiment of the invention, approximately 0.01 to 2000 μg and, preferably, 0.1 to 500 μg, of the PMP protein or PMP-derived protein is administered is to a host. Compositions which induce T-cell responses which are bactericidal or reactive with host cells infected with *Chlamydia* are also an aspect of the invention. Preferred are compositions additionally comprising an adjuvant.

The combined immunogen and carrier or diluent may be an aqueous solution, emulsion or suspension or may be a dried preparation. Appropriate carriers are known to those skilled in the art and include stabilizers, diluents, and buffers. Suitable stabilizers include carbohydrates, such as sorbitol, lactose, mannitol, starch, sucrose, dextran, and glucose, and proteins, such as albumin or casein. Suitable diluents include saline, Hanks Balanced Salts, and Ringers solution. Suitable buffers include an alkali metal phosphate, an alkali metal carbonate, or an alkaline earth metal carbonate. In preferred embodiments, the composition of the invention is formulated for administration to humans.

The immunogenic compositions, including vaccines, of the invention are prepared by techniques known to those skilled in the art, given the teachings contained herein. Generally, an immunogen is mixed with the carrier to form a solution, suspension, or emulsion. One or more of the additives discussed herein may be added in the carrier or may be added subsequently. The vaccine preparations may be desiccated or lyophilized, for example, by freeze drying or spray drying for storage or formulations purposes. They may be subsequently reconstituted into liquid vaccines by the addition of an appropriate liquid carrier or administered in dry formulation using methods known to those skilled in the art, particularly in capsules or tablet forms.

An effective amount of the antigenic, immunogenic, pharmaceutical, including, but not limited to vaccine, composition of the invention should be administered, in which "effective amount" is defined as an amount that is sufficient to produce a desired prophylactic, therapeutic or ameliorative response in a subject, including but not limited to an immune response. The amount needed will vary depending upon the immunogenicity of

the PMP protein, PMP-derived polypeptide or nucleic acid used, and the species and weight of the subject to be administered, but may be ascertained using standard techniques.

Immunogenic, antigenic, pharmaceutical and vaccine compositions may further contain one or more auxiliary substance, such as wetting or emulsifying agents, pH buffering agents, or adjuvants to enhance the effectiveness thereof. Immunogenic, antigenic, pharmaceutical and vaccine compositions may be administered to birds, humans or other mammals, including ruminants, rodents or primates, by a variety of administration routes, including parenterally, intradermally, intraperitoneally, subcutaneously or intramuscularly.

10 Alternatively, the immunogenic, antigenic, pharmaceutical and vaccine compositions formed according to the present invention, may be formulated and delivered in a manner to evoke an immune response at mucosal surfaces. Thus, the immunogenic, antigenic, pharmaceutical and vaccine compositions may be administered to mucosal surfaces by, for example, the nasal, oral (intragastric), ocular, bronchiolar, intravaginal or 15 intrarectal routes. Alternatively, other modes of administration including suppositories and oral formulations may be desirable. For suppositories, binders and carriers may include, for example, polyalkalene glycols or triglycerides. Oral formulations may include normally employed incipients such as, for example, pharmaceutical grades of saccharine, cellulose and magnesium carbonate. These compositions can take the form of microspheres, 20 solutions, suspensions, tablets, pills, capsules, sustained release formulations or powders and contain about 0.001 to 95% of the PMP protein. The immunogenic, antigenic, pharmaceutical and vaccine compositions are administered in a manner compatible with the dosage formulation, and in such amount as will be therapeutically effective, protective or immunogenic. Preferred are compositions additionally comprising an adjuvant.

Further, the immunogenic, antigenic, pharmaceutical and vaccine compositions may be used in combination with or conjugated to one or more targeting molecules for delivery to specific cells of the immune system and/or mucosal surfaces. Some examples include but are not limited to vitamin B12, bacterial toxins or fragments thereof, monoclonal antibodies and other specific targeting lipids, proteins, nucleic acids or carbohydrates.

Suitable regimes for initial administration and booster doses are also variable, but may include an initial administration followed by subsequent administrations. The dose may also depend on the route(s) of administration and will vary according to the size of the host. The concentration of the PMP protein or PMP-derived polypeptide in an

antigenic, immunogenic or pharmaceutical composition according to the invention is in general about 0.001 to 95%.

The antigenic, immunogenic or pharmaceutical preparations, including vaccines, may comprise as the immunostimulating material a nucleic acid vector comprising at least a portion of the nucleic acid molecule encoding the PMP protein or PMP-derived polypeptide.

A vaccine comprising nucleic acid molecules encoding one or more PMP polypeptides, PMP-derived polypeptides or fusion proteins as described herein, such that the polypeptide is generated in situ is provided. In such vaccines, the nucleic acid 10 molecules may be present within any of a variety of delivery systems known to those skilled in the art, including nucleic acid expression systems, bacterial and viral expression systems. Appropriate nucleic acid expression systems contain the necessary nucleotide sequences for expression in the patient such as suitable promoter and terminating signals. In a preferred embodiment, the nucleic acid molecules may be introduced using a viral expression system 15 (e.g., vaccinia or other pox virus, alphavirus retrovirus or adenovirus) which may involve the use of non-pathogenic (defective) virus. Techniques for incorporating nucleic acid molecules into such expression systems are well known to those of ordinary skill in the art. The nucleic acid molecules may also be administered as "naked" plasmid vectors as described, for example, in Ulmer et al. Science 259:1745-1749 (1992) and reviewed by 20 Cohen, Science 259:1691-1692 (1993). Techniques for incorporating DNA into such vectors are well known to those of ordinary skill in the art. A retroviral vector may additionally transfer or incorporate a gene for a selectable marker (to aid in the identification or selection of transduced cells) and/or a targeting moiety, such as a gene that encodes a ligand for a receptor on a specific target cell, to render the vector target specific. 25 Targeting may also be accomplished using an antibody, by methods know to those skilled in

Nucleic acid molecules (DNA or RNA) of the invention can be administered as vaccines for therapeutic or prophylactic purpose. Typically a DNA molecule is placed under the control of a promoter suitable for expression in a mammalian cell. The promoter can function ubiquitously or tissue-specifically. Examples of non-tissue specific promoters include the early cytomegalovirus (CMV) promoter (described in U.S. Patent No. 4,168,062) and Rous Sarcoma virus promoter (described in Norton and Coffin, Molec. Cell Biol. 5:281 (1985)). The desmin promoter (Li et al. Gene 78:243 (1989); Li & Paulin, J. Biol Chem 266:6562 (1991); and Li & Paulin, J. Biol Chem 268:10401 (1993)) is tissue

the art.

specific and drives expression in muscle cells. More generally, useful vectors are described in, e.g., WO 94/21797 and Hartikka et al., <u>Human Gene Therapy</u> 7:1205 (1996).

A composition of the invention can contain one or several nucleic acid molecules of the invention. It can also contain at least one additional nucleic acid molecule encoding another antigen or fragment derivative, including but not limited to, DPT vaccines, HMWP of *C. trachomatis*, MOMP of *C. trachomatis*, entire organisms or subunits therefrom of Chlamydia, Neisseria, HIV, *Haemophilus influenzae*, *Moraxella catarrhalis*, *Human papilloma virus*, *Herpes simplex virus*, *Haemophilus ducreyi*, *Treponema pallidium*, *Candida albicans* and *Streptococcus pneumoniae*, etc. A nucleic acid molecule encoding a cytokine, such as interleukin-1 or interleukin-12 can also be added to the composition so that the immune response is enhanced. DNA molecules of the invention and/or additional DNA molecules may be on different plasmids or vectors in the same composition or can be carried in the same plasmid or vector.

Other formulations of nucleic acid molecules for therapeutic and
prophylactic purposes include sterile saline or sterile buffered saline colloidal dispersion systems, such as macromolecule complexes, nanocapsules, silica microparticles, tungsten microparticles, gold microparticles, microspheres, beads and lipid based systems including oil-in-water emulsions, micelles, mixed micelles and liposomes. A preferred colloidal system for use as a delivery vehicle *in vitro* and *in vivo* is a liposome (*i.e.*, an artificial vesicle). The uptake of naked nucleic acid molecules may be increased by incorporating the nucleic acid molecules into and/or onto biodegradable beads, which are efficiently transported into the cells. The preparation and use of such systems is well known in the art.

A nucleic acid molecule can be associated with agents that assist in cellular uptake. It can be formulated with a chemical agent that modifies the cellular permeability, such as bupivacaine (see, *e.g.*, WO 94/16737).

Cationic lipids are also known in the art and are commonly used for DNA delivery. Such lipids include Lipofectin[™], also known as DOTMA (N-[1-(2,3-dioleyloxy)propyl]-N,N,N-trimethylammonium chloride), DOTAP (1,2-bis(oleyloxy)-3-(trimethylammonio)propane, DDAB (dimethyldioctadecylammonium bromide), DOGS (dioctadecylamidologlycy spermine) and cholesterol derivatives such as DC-Chol (3 beta-(N-(N',N'-dimethyl aminomethane)-carbamoyl) cholesterol. A description of these cationic lipids can be found in EP 187,702, WO 90/11092, U.S. Patent No. 5,283,185, WO 91/15501, WO 95/26356, and U.S. Patent No. 5,527,928. Cationic lipids for DNA delivery are preferably used in association with a neutral lipid such as DOPE (dioleyl

35 phosphatidylethanolamine) as described in, e.g., WO 90/11092.

Other transfection facilitation compounds can be added to a formulation containing cationic liposomes. They include, *e.g.*, spermine derivatives useful for facilitating the transport of DNA through the nuclear membrane (see, for example, WO 93/18759)and membrane-permeabilizing compounds such as GALA, Gramicidine S and cationic bile salts (see, for example, WO 93/19768).

The amount of nucleic acid molecule to be used in a vaccine recipient depends, *e.g.*, on the strength of the promoter used in the DNA construct, the immunogenicity of the expressed gene product, the mode of administration and type of formulation. In general, a therapeutically or prophylactically effective dose from about 1 μg to about 1 mg, preferably from about 10 μg to about 800 μg and more preferably from about 25 μg to about 250 μg can be administered to human adults. The administration can be achieved in a single dose or repeated at intervals.

The route of administration can be any conventional route used in the vaccine field. As general guidance, a nucleic acid molecule of the invention can be

15 administered via a mucosal surface, *e.g.*, an ocular, intranasal, pulmonary, oral, intestinal, rectal, vaginal, and urinary tract surface; or via a parenteral route, *e.g.*, by an intravenous, subcutaneous, intraperitoneal, intradermal, intra-epidermal or intramuscular route. The choice of administration will depend on the formulation that is selected. For instance a nucleic acid molecule formulated in association with bupivacaine is advantageously

20 administered into muscles.

Recombinant bacterial vaccines genetically engineered for recombinant expression of nucleic acid molecules encoding PMP or PMP-derived polypeptides include *Shigella, Salmonella, Vibrio cholerae*, and *Lactobacillus*. Recombinant BCG and *Streptococcus* expressing PMP or PMP-derived polypeptides can also be used for prevention or treatment of *Chlamydia* infections.

Non-toxicogenic *Vibrio cholerae* mutant strains that are useful as a live oral vaccine are described in Mekalanos et al. Nature 306:551 (1983) and U.S. Patent No. 4,882,278. An effective vaccine dose of a *Vibrio cholerae* strain capable of expressing a polypeptide or polypeptide derivative encoded by a DNA molecule of the invention can be administered. Preferred routes of administration include all mucosal routes, most preferably intranasally or orally.

Attenuated *Salmonella typhimurium* strains, genetically engineered for recombinant expression of heterologous antigens or not and their use as oral vaccines are described in Nakayama et al. <u>Bio/Technology</u> 6:693 (1988) and WO 92/11361. Preferred routes of administration include all mucosal routes, most preferably intranasally or orally.

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Other bacterial strains useful as vaccine vectors are described in High et al., <u>EMBO</u> 11:1991(1992); Sizemore et al., <u>Science</u> 270:299 (1995) (*Shigella flexneri*); Medaglini et al., <u>Proc Natl. Acad. Sci. US</u> 92:6868 (1995) (*Streptococcus gordonii*); and Flynn, <u>Cell Mol. Biol</u>.40:31 (1994); WO 88/6626; WO 90/0594; WO 91/13157; WO 92/1796; and WO 02/21376 (*Bacille Calmette Guerin*).

In genetically engineered recombinant bacterial vectors, nucleic acid molecule(s) of the invention can be inserted into the bacterial genome, carried on a plasmid, or can remain in a free state.

When used as vaccine agents, recombinant bacterial vaccines, nucleic acid molecules and polypeptides of the invention can be used sequentially or concomitantly as part of a multistep immunization process. For example, a mammal or bird can be initially primed with a vaccine vector of the invention such as pox virus, *e.g.*, via the parenteral route and then boosted several time with the a polypeptide *e.g.*, via the mucosal route. In another example, a mammal can be vaccinated with polypeptide via the mucosal route and at the same time or shortly thereafter, with a nucleic acid molecule via intramuscular route.

An adjuvant can also be added to a composition containing a vaccine recombinant bacteria. To efficiently induce humoral immune responses (HIR) and cell-mediated immunity (CMI), immunogens are typically emulsified in adjuvants.

Immunogenicity can be significantly improved if the immunogen is co-administered with an adjuvant. Adjuvants may act by retaining the immunogen locally near the site of administration to produce a depot effect facilitating a slow, sustained release of antigen to cells of the immune system. Adjuvants can also attract cells of the immune system to an immunogen depot and stimulate such cells to elicit immune responses.

Many adjuvants are toxic, inducing granulomas, acute and chronic

25 inflammations (Freund's complete adjuvant, FCA), cytolysis (saponins and Pluronic polymers) and pyrogenicity, arthritis and anterior uveitis (LPS and MDP). Although FCA is an excellent adjuvant and widely used in research, it is not licensed for use in human or veterinary vaccines because of its toxicity.

Immunostimulatory agents or adjuvants have been used for many years to improve the host immune responses to, for example, vaccines. Intrinsic adjuvants, such as lipopolysaccharides, normally are the components of the killed or attenuated bacteria used as vaccines. Extrinsic adjuvants are immunomodulators which are typically non-covalently linked to antigens and are formulated to enhance the host immune responses. Thus, adjuvants have been identified that enhance the immune response to antigens delivered parenterally. Aluminum hydroxide, aluminum oxide, and aluminum phosphate (collectively

commonly referred to as alum) are routinely used as adjuvants in human and veterinary vaccines. The efficacy of alum in increasing antibody responses to diphtheria and tetanus toxoids is well established and a HBsAg vaccine has been adjuvanted with alum.

Other extrinsic adjuvants may include chemokines, cytokines (e.g., IL-2), saponins complexed to membrane protein antigens (immune stimulating complexes), pluronic polymers with mineral oil, killed mycobacteria in mineral oil, Freund's complete adjuvant, bacterial products, such as muramyl dipeptide (MDP) and lipopolysaccharide (LPS), as well as lipid A, and liposomes.

International Patent Application PCT/US95/09005, incorporated herein by reference, describes mutated forms of heat labile toxin of enterotoxigenic *E. coli* ("mLT"). U.S. Patent No. 5,057,540, incorporated herein by reference, describes the adjuvant, QS21, an HPLC purified non-toxic fraction of a saponin from the bark of the South American tree *Quiliaja saponaria molina*. 3D-MPL is described in Great Britain Patent 2,220,211, which is incorporated herein by reference.

U.S. Patent No. 4,855,283 granted to Lockhoff et al. on August 8, 1989, which is incorporated herein by reference, teaches glycolipid analogues including N-glycosylamides, N-glycosylureas and N-glycosylcarbamates, each of which is substituted in the sugar residue by an amino acid, as immuno-modulators or adjuvants. Lockhoff reported that N-glycosphospholipids and glycoglycerolipids are capable of eliciting strong immune responses in both herpes simplex virus vaccine and pseudorabies virus vaccine. Some glycolipids have been synthesized from long chain-alkylamines and fatty acids that are linked directly with the sugars through the anomeric carbon atom, to mimic the functions of the naturally occurring lipid residues.

U.S. Patent No. 4,258,029 granted to Moloney, incorporated herein by reference, teaches that octadecyl tyrosine hydrochloride (OTH) functions as an adjuvant when complexed with tetanus toxoid and formalin inactivated type I, II and III poliomyelitis virus vaccine. Lipidation of synthetic peptides has also been used to increase their immunogenicity.

Therefore, according to the invention, the immunogenic, antigenic,
30 pharmaceutical, including vaccine, compositions comprising a PMP protein, or a PMP
derived polypeptide or a PMP encoding nucleic acid or fragment thereof, vector or cell
expressing the same, may further comprise an adjuvant, such as, but not limited to alum,
mLT, QS21, MMPL, CpG DNA, MF59, calcium phosphate, PLG and all those listed above.
Preferably, the adjuvant is selected from one or more of the following: alum, QS21, CpG
35 DNA, PLG, LT, 3D-mPL, or Bacille Calmette-Guerine (BCG) and mutated or modified

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forms of the above, particularly mLT and LTR192G. The compositions of the present invention may also further comprise a suitable pharmaceutical carrier, including but not limited to saline, bicarbonate, dextrose or other aqueous solution. Other suitable pharmaceutical carriers are described in Remington's Pharmaceutical Sciences, Mack Publishing Company, a standard reference text in this field, which is incorporated herein by reference in its entirety.

Immunogenic, antigenic and pharmaceutical, including vaccine, compositions may be administered in a suitable, nontoxic pharmaceutical carrier, may be comprised in microcapsules, and/or may be comprised in a sustained release implant.

Immunogenic, antigenic and pharmaceutical, including vaccine, compositions may desirably be administered at several intervals in order to sustain antibody levels and/or T-cell levels. Immunogenic, antigenic and pharmaceutical, including vaccine, compositions may be used in conjunction with other bacteriocidal or bacteriostatic methods.

Another embodiment of the vaccines of the present is a vaccine comprising
one or more isolated PMPE or PMPI polypeptides of *Chlamydia spp*, having a molecular weight between 90 and 115 kDa as determined in SDS polyacrylamide gel electrophoresis; or isolated nucleic acids encoding an isolated PMPE or PMPI polypeptide of *Chlamydia spp*., having a molecular weight between 90 and 115 kDa as determined in SDS polyacrylamide gel electrophoresis, and further comprising one or more components selected from the group consisting of alum, MLT, QS21, MF59, CpG DNA, MPL, calcium phosphate and PLG.

Also included in the invention is a method of producing an immune response in an animal comprising immunizing the animal with an effective amount of one or more of the PMP polypeptides or nucleic acid molecules encoding the PMP polypeptides of the invention, compositions comprising the same and vaccines comprising the same. The PMP polypeptides, nucleic acids, compositions and vaccines comprising the PMP polypeptides of the invention may be administered simultaneously or sequentially. Examples of simultaneous administration include cases in which two or more polypeptides, nucleic acids, compositions, or vaccines, which may be the same or different, are administered in the same or different formulation or are administered separately, *e.g.*, in a different or the same formulation but within a short time (such as minutes or hours) of each other. Examples of sequential administration include cases in which two or more polypeptides, nucleic acids, compositions or vaccines, which may be the same or different, are not administered together or within a short time of each other, but may be administered separately at intervals of, for example, days, weeks, months or years.

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The polypeptides, nucleic acid molecules or recombinant bacterial vaccines of the present invention are also useful in the generation of antibodies, as described *supra*, or T-cells. For T-cells, animals, including humans, are immunized as described above. Following immunization, PBL (peripheral blood lymphocytes), spleen cells or lymph node cells are harvested and stimulated *in vitro* by placing large numbers of lymphocytes in flasks with media containing human serum. A polypeptide of the present invention is added to the flasks, and T-cells are harvested and placed in new flasks with X-irradiated peripheral blood mononuclear cells. The polypeptide is added directly to these flasks, and cells are grown in the presence of IL-2. As soon as the cells are shown to be *Chlamydia* specific T-cells, they are changed to a stimulation cycle with higher IL-2 concentrations (20 units) to expand them.

Alternatively, one or more T-cells that proliferate in the presence of a polypeptide of the present invention can be expanded in number by cloning. Methods for cloning cells are well known in the art. For example, T-cell lines may be 15 established in vitro and cloned by limiting dilution. Responder T-cells are purified from the peripheral blood established in culture by stimulating with the nominal antigen in the presence of irradiated autologous filler cells. In order to generate CD4⁺ T-cell lines, the Chlamydia polypeptide is used as the antigenic stimulus and autologous PBL or lymphoblastoid cell lines (LCL) immortalized by infection with Epstein Barr virus are used 20 as antigen presenting cells. In order to generate CD8+ T-cell lines, autologous antigenpresenting cells transfected with an expression vector which produces the relevant Chlamydia polypeptide may be used as stimulator cells. T-cell lines are established following antigen stimulation by plating stimulated T-cells in 96-well flat-bottom plates with PBL or LCL cells and recombinant interleukin-2 (rIL2) (50 U/ml). Wells with 25 established clonal growth are identified at approximately 2-3 weeks after initial plating and restimulated with appropriate antigen in the presence of autologous antigen-presenting cells, then subsequently expanded by the addition of low doses of IL2. T-cell clones are maintained in 24-well plates by periodic restimulation with antigen and IL2 approximately every two weeks.

T-cell preparations may be further enriched by isolating T-cells specific for antigen reactivity using the methods disclosed by Kendricks et al. in U.S. Patent No. 5,595,881.

The vaccine compositions of the present inventions are useful in preventing, treating or ameliorating disease symptoms in an animal with a disease or disorder associated with *Chlamydia* infection. Such diseases or disorders include, but are not limited to,

Chlamydia bacterial infection, conjunctivitis, urethritis, lymphogranuloma venereum (LGV), cervicitis, epididymitis, salpingitis, tubal occlusion, infertility, cervical cancer, reactive arthritis, arteriosclerosis and atherosclerosis.

5 5.8 IMMUNOASSAYS AND DIAGNOSTIC REAGENTS

The PMP protein, PMP-derived polypeptide or nucleic acid encoding same, and fragments thereof, are useful as diagnostic reagents. Use of the proteins and nucleic acids of the invention as an antigen or immunogen for the generation of anti-PMP protein antibodies or as an antigen in immunoassays including enzyme-linked immunosorbent assays (ELISA), radioimmmunoassays (RIA) and other non-enzyme linked antibody binding assays or procedures known in the art for the detection of anti-bacterial, anti-*Chlamydia*, and anti-PMP protein antibodies are encompassed by the invention.

ELISA is well-known and routine in the art. Generally, in ELISA, the PMP protein is immobilized onto a selected surface, for example, a surface capable of binding proteins such as the wells of a polystyrene microtiter plate. After washing to remove incompletely absorbed PMP protein, a nonspecific protein solution that is known to be antigenically neutral with regard to the test sample may be bound to the selected surface. This allows for blocking of nonspecific absorption sites on the immobilizing surface and thus reduces the background caused by nonspecific bindings of antisera onto the surface.

The immobilizing surface is then contacted with a sample, such as clinical or biological materials, to be tested in a manner conducive to immune complex (antigen/antibody) formation. This may include diluting the sample with diluents, such as solutions of bovine gamma globulin (BGG) and/or phosphate buffered saline (PBS)/Tween. The sample is then allowed to incubate for from 2 to 4 hours, at temperatures on the order of about 20°C to 37°C. Following incubation, the sample-contacted surface is washed to remove non-immunocomplexed material. The washing procedure may include washing with a solution, such as PBS/Tween or a borate buffer. Following formation of specific immunocomplexes between the test sample and the bound PMP protein, and subsequent washing, the occurrence, and even amount, of immunocomplex formation may be determined by subjecting the immunocomplex to a second antibody having specificity for the first antibody. If the test sample is of human origin, the second antibody is an antibody having specificity for human immunoglobulins and in general IgG.

To provide detecting means, the second antibody may have an associated activity such as an enzymatic activity that will generate, for example, a color development upon incubating with an appropriate chromogenic substrate. Detection may then be

achieved by detecting color generation. Quantification may then be achieved by measuring the degree of color generation using, for example, a visible spectrophotometer and comparing to an appropriate standard. Any other detecting means known to those skilled in the art are included.

In Western blot assays, the polypeptide, either as a purified preparation or a cell extract, is subjected to SDS-PAGE electrophoresis, for example, as described by Laemmli, Nature 227:690 (1970) or any other method known in the art. After transfer to a nitrocellulose membrane, the material is further incubated with the serum sample, polyclonal antibody preparation, or monoclonal antibody diluted in the range of from about 1:5 to 1:5000, preferably from about 1:100 to about 1:500, depending upon the titer and specification of the antibodies. Appropriate dilutions can be readily determined by methods known in the art. The reaction is revealed according to standard procedures. For example, when human antibody is used, the membrane is incubated in a goat anti-human peroxidase conjugate for an appropriate length of time. The membrane is washed. The reaction is developed with the appropriate substrate and stopped. The reaction is measured visually by the appearance of a colored band *e.g.*, by colorimetry.

In a dot blot assay, the purified or partially purified polypeptide or cell extract can be used. Briefly, a solution of the antigen at about 100 μg/ml is serially two-fold diluted in 50m M Tris-HCl (pH 7.5). 100 μl of each dilution is applied to a 0.45 μm nitrocellulose membrane set in a 96-well dot blot apparatus. The buffer is removed by applying vacuum to the system. Wells are washed by addition of 50μM Tris-HCl (pH 7.5) and the membrane is air-dried. The membrane is saturated in block buffer (50m M Tris-HCl (pH 7.5), 0.15 M NaCl and 10 g/L skim milk) and incubated with an antiserum dilution from about 1:50 to about 1:500. The reaction is revealed according to standard procedures.

25 For example, a goat anti-rabbit peroxidase conjugate is added to the well when rabbit antibodies are used. Incubation is carried out 90 minutes at 37°C and the blot is washed. The reaction is developed with the appropriate substrate and stopped. The reaction is measured visually by the appearance of a colored spot, *e.g.*, by colorimetry.

The PMP proteins, PMP-derived polypeptides or nucleic acids encoding same, and fragments thereof, are also useful as antigens or immunogens for the generation of anti-PMP protein T-cell response or as an antigen in immunoassays, including T-cell proliferation assays, cytokine production, delayed hypersensitivity reactions or cytotoxic T-cells (CTL) reactions.

For analysis of *Chlamydia* peptide specific T-cell proliferative responses, 35 fresh peripheral blood, spleen or lymph node cells are harvested. Cells are plated into 96well round bottom microtiter plates and are incubated with peptides. Data is expressed as a stimulation index (SI) which is defined as the mean of the number of cells in experimental wells divided by the mean of the number of cells in control wells (no antigen).

For analysis of cytokine release of T-cells in response to *Chlamydia*5 polypeptides, responder cells are mixed with polypeptides. Supernatants are collected and added to an ELISA coated with antibody to the cytokine (*e.g.*, anti-IFN-α or anti-IL-2 antibody). After washing, rabbit anti-cytokine polyclonal antibody (*e.g.*, anti- IFN-α or anti-IL-2) is added. Labeled goat anti-rabbit IgG polyclonal is added. Substrate is added and the amount of cytokine released into the supernatant is determined based upon the amount of color developed in the ELISA.

Another embodiment includes diagnostic kits comprising all of the essential reagents required to perform a desired immunoassay according to the present invention. The diagnostic kit may be presented in a commercially packaged form as a combination of one or more containers holding the necessary reagents. Such a kit may comprise PMP protein, PMP-derived polypeptide or nucleic acid encoding same, or a monoclonal or polyclonal antibody of the present invention, in combination with several conventional kit components. Conventional kit components will be readily apparent to those skilled in the art and are disclosed in numerous publications, including, for example, Harlow and Lane, Antibodies A Laboratory Manual, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York 1988) which is incorporated herein by reference in its entirety. Conventional kit components may include such items as, for example, microtiter plates, buffers to maintain the pH of the assay mixture (such as, but not limited to Tris, HEPES, etc.), conjugated second antibodies, such as peroxidase conjugated anti-mouse IgG (or any anti-IgG to the animal from which the first antibody was derived) and the like, and other standard reagents.

The nucleic acid molecules containing the PMP encoding sequences of the present invention may be used in combination with an appropriate indicator means, such as a label, for determining hybridization. A wide variety of appropriate indicator means are known in the art, including radioactive, enzymatic or other ligands, such as avidin/biotin and digoxigenin-labeling, which are capable of providing a detectable signal. In some diagnostic embodiments, an enzyme tag, such as urease, alkaline phosphatase or peroxidase, instead of a radioactive tag, may be used. In the case of enzyme tags, colorimetric indicator substrates are known which can be employed to provide a means visible to the human eye or spectrophotometrically, to identify specific hybridization with samples containing PMP

35 protein gene sequences.

Probes of the invention can be used in diagnostic tests, as capture or detection probes. Such capture probes can be conventionally immobilized on a solid support directly or indirectly, by covalent means or by passive adsorption. A detection probe can be labeled by a detection marker selected from radioactive isotopes, enzymes, such as peroxidase, alkaline phosphatase, and enzymes able to hydrolyze a chromogenic, fluorogenic or luminescent substrate; compounds that are chromogenic, fluorogenic or luminescent; nucleotide base analogs; and biotin.

Probes of the invention can be used in any conventional hybridization techniques, such as dot blot (Maniatis et al., Molecular Cloning: A Laboratory Manual (1982) Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York 1982), Southern blot (Southern, J. Mol. Biol. 98:503 1975), northern blot (identical to Southern blot to the exception that RNA is used as a target), or sandwich techniques (Dunn et al., Cell 12:23 1977).

In embodiments involving solid-phase procedures, the test DNA (or RNA)

from samples, such as clinical samples, including exudates, body fluids (*e.g.*, serum, amniotic fluid, middle ear effusion, sputum, semen, urine, tears, mucus, bronchoalveolar lavage fluid) or even tissues, is absorbed or otherwise affixed to a selected matrix or surface. The fixed, single-stranded nucleic acid is then subjected to specific hybridization with selected probes comprising the nucleotide sequences encoding a PMP protein, or fragments or analogues thereof, under desired conditions. The selected conditions will depend on the particular criteria required and on, for example, the G+C content, type of target nucleic acid, source of nucleic acid, size of hybridization probe, etc. Following washing of the hybridization surface so as to remove non-specifically bound probe molecules, specific hybridization is detected, or even quantified, by means of the label. It is preferred to select nucleotide acid sequence portions that are conserved among species of *Chlamydia*. The selected probe may be at least 15 bp and may be in the range of about 30 to 90 bp.

The invention also relates to methods for identifying compounds which interact with and inhibit or activate an activity of the polypeptides or nucleic acid molecules of the invention comprising contacting a composition comprising the polypeptide or the nucleic acid molecule with the compound to be screened under conditions that permit interaction between the compound and the polypeptide or nucleic acid molecule to assess the interaction of a compound and to detect interaction of the compound with the polypeptide of nucleic acid. The interaction of the compound with the polypeptide or nucleic acid molecule is determined by the association of a second component (*e.g.*, an

antibody) capable of providing a detectable signal in response to the interaction of the polypeptide or nucleic acid molecule with the compound; and determining the presence or absence of a signal generated from the interaction of the compound with the polypeptide or nucleic acid molecule. Alternatively, the interaction of the compound with the polypeptide or nucleic acid molecule is determined by the ability of the compound to inhibit the activity of the polypeptide or the nucleic acid molecule.

5.9 APPLICATIONS

The proteins, polypeptides, peptides, antibodies, T-cells and nucleic acids of the invention are useful as reagents for clinical or medical diagnosis of *Chlamydia* infections and for scientific research on the properties of pathogenicity, virulence, and infectivity of *Chlamydia*, as well as host defense mechanisms. For example, DNA and RNA of the invention can be used as probes to identify the presence of *Chlamydia* in biological specimens by hybridization or PCR amplification. The DNA and RNA can also be used to identify other bacteria that might encode a polypeptide related to the *Chlamydia* PMP protein. The proteins of the invention may be used to prepare polyclonal and monoclonal antibodies that can be used to further purify compositions containing the proteins of the invention by affinity chromatography or for use as diagnostic or as prophylactic or therapeutic agents. The proteins can also be used in standard immunoassays to screen for the presence of antibodies or T-cells to *Chlamydia* in a biological sample.

5.10. BIOLOGICAL DEPOSITS

Certain plasmids that contain portions of the gene having the open reading
frame of the PMP genes encoding the PMP proteins of the present invention have been inserted into *E. coli* and deposited with the American Type Culture Collection (ATCC) located at 10801 University Boulevard, Manassas, Virginia 20110-2209, U.S.A., pursuant to the Budapest Treaty and pursuant to 37 CFR 1.808 and prior to the filing of this application. The identifications of the respective portions of the genes present in these plasmids are shown below.

Samples of the deposited materials will become available to the public upon grant of a patent based upon this United Stated patent application. The invention described and claimed herein is not to be limited by the scope of the plasmids deposited, since the deposited embodiment is intended only as an illustration of the invention. Any equivalent

or similar plasmids that encode similar or equivalent proteins or fragments or analogues thereof as described in this application are within the scope of the invention.

	Plasmid	ATCC Accession No.	Date Deposited
5	M15 pREP (pQE-pmpE)#37	ATCC PTA-2462	Sept. 12, 2000
	TOP10(pBAD-pmpI-Ct-Uni)#7	ATCC PTA-2461	Sept. 12, 2000

The above disclosure generally describes the present invention. A more complete understanding can be obtained by reference to the following specific examples.

These examples are described solely for purposes of illustration and not intended to limit the scope of the invention. Changes in form and substitution of equivalents are contemplated as circumstances may suggest or render expedient. Although specific terms have been employed herein such terms are intended in a descriptive sense and not for purposes of limitation.

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6. EXAMPLES

The above disclosure generally describes the present invention. The examples are described solely for the purpose of illustration and are not intended to limit the scope of the invention. Changes in form and substitution of equivalents are contemplated as circumstances suggest or render expedient. Although specific terms have been employed herein, such terms are intended in a descriptive sense and not for purposes of limitation.

Methods of molecular genetics, protein biochemistry and immunology used but not explicitly described in the disclosure and examples are amply reported in the scientific literature and are well within the ability of those skilled in the art.

25

6.1 EXTRACTION OF ENVELOPE PROTEINS

McCoy cells are cultured either in standard 225 cm² tissue culture flasks or in Bellco spinner flasks (Cytodex microcarrier, Pharmacia) at 37°C in 5% CO₂ using DMEM media supplemented with 10% *Chlamydia*-antibody-free fetal bovine serum, glucose and nonessential amino acids. *C. trachomatis*, including but not limited to the L2 serovar (ATCC VR-902B), elementary bodies (EBs) are prepared from lysates of infected McCoy cells. Basically, McCoy cells infected with *C. trachomatis* are sonicated and cellular debris is removed by centrifugation. The supernatant containing Chlamydial EBs is then centrifuged and the pellet containing the EBs is resuspended in Hanks' balanced salts solution (HBSS). RNAase/DNAase solution is added and incubated at 37°C for 1 hour

with occasional mixing. The EB containing solution is layered onto a discontinuous density gradient (40%, 44% and 54%) of Renogratin-60 (mixture of diatrizoate melgumine and diatrizoate sodium, Bracco Diagnostics, Princeton, NJ) and ultracentrifuged for separation of the EBs on the gradient. After centrifugation, the EBs are harvested from the gradient between the interface of the 44% and 54% layers. The EBs are washed in phosphate buffered saline and resuspended in HBSS.

Purified EBs are sequentially extracted with 0.1% OGP [high ionic strength] in HBSS to remove peripheral surface proteins and held on ice. The same EB preparation is then extracted with 1.0% OGP, 10 mM DTT, 1 mM PMSF, and 10 mM EDTA, in a 50 mM 10 Tris pH 7.4 buffer. The extracts are dialyzed (3500 molecular weight cut off) to remove detergent and other reagents and are concentrated by lyophilization. Eluents are dialyzed to remove excess detergent and then lyophilized. Envelope proteins are size fractionated by SDS-PAGE and visualized by silver staining or analyzed by Western blotting. Protein(s) of about 90-115 kDa present in moderate amounts are detected in the gel.

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6.2 AMINO TERMINAL SEQUENCING OF PMP POLYPEPTIDE

To obtain the N-terminal amino acid sequence, sufficient quantities of the PMPE or PMPI protein (≥ 5 μg) are electroblotted onto a PVDF membrane (Applied Biosystems), and stained with Coomassie blue. Immobilized protein is released from the membrane and treated *in situ* with low levels of endopeptidase Lys-C, endopeptidase Arg-C and/or endopeptidase Glu-C to fragment the native protein. The resulting peptide fragments are purified by HPLC and their N-terminal amino acid sequences are determined using an ABI 430 Protein Sequenator and standard protein sequencing methodologies.

25 6.3. DETERMINATION OF POST-TRANSLATIONAL MODIFICATIONS

Recently, several *C. trachomatis* membrane-associated proteins have been shown to be post-translationally modified. The 18 kDa and 32 kDa cysteine-rich EB proteins, which are lectin-binding proteins, have been shown to carry specific carbohydrate moieties (Swanson et al. 1990. <u>Infect. Immun.</u> 58:502-507). Incorporation of radiolabeled palmitic acid has been used to demonstrate that the about 27 kDa *C. trachomatis* Mip-like protein is lipidated (Lundemose et al. 1993. <u>J. Bacteriol.</u> 175:3669-3671). Swanson et al. have discovered that the MOMP from the L2 serovar contains N-acetylglucosamine and/or N-acetylgalactosamine and these carbohydrate moieties mediate binding of MOMP to Hela cell membranes.

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To ascertain whether the PMPE or PMPI protein is glycosylated, EBs are grown on McCoy cells in the presence of tritiated galactose or glucosamine and analyzed by SDS-PAGE and autoradiography. Briefly, McCoy cells are grown in T225 flasks under standard conditions (DMEM + 10% FCS, 35 ml per flask, 10% CO₂) to about 90% confluency and inoculated with sufficient EBs to achieve 90%-100% infectivity. Following a 3 hour infection period at 37°C cycloheximide is added (1 μg/ml) to inhibit host cell protein synthesis and the cultures reincubated for an additional 4-6 hours. Approximately 0.5 mCi of tritiated galactose (D-[4,5-³H(N)]galactose, NEN) or glucosamine (D- [1,6-³H(N)glucosamine, NEN) is then be added to each flask and the cultures allowed to incubate for an additional 30-40 hours. Cells are harvested by scraping, and EBs purified by gradient centrifugation. PMPE or PMPI protein is isolated from 1.0% OGP surface extracts, eluted with NaCl and analyzed by SDS-PAGE using ¹⁴C-labeled molecular weight markers (BRL). The resulting gel is dried and subjected to autoradiography by exposure for 1-4 weeks to Kodak X-AR film at -70°C.

To determine post synthesis lipid modification, *C. trachomatis* is cultivated on monolayers of McCoy cells according to standard procedures. Approximately 24 hours postinfection, conventional culture media (DMEM + 10% FCS) is removed and replaced with a serum-free medium containing cycloheximide (1 μg/ml) and [U-¹⁴C]palmitic acid (0.5 mCi/T225 flask, NEN) and incubated for a further 16-24 hours to allow protein lipidation to occur. Surface EB extracts are prepared and analyzed by autoradiography as described above.

6.4 ANTI-PMPE or Anti-PMPI ANTISERUM

Antisera to PMPE or PMPI polypeptides are prepared by injecting the PMPE or PMPI polypeptide into an animal, such as a rabbit, mouse or guinea pig, with or without an adjuvant by any method generally known to those skilled in the art. For instance, PMPE is injected with Freund's complete adjuvant followed by injections of PMPE with Freund's incomplete adjuvant. Normally, a semi-purified or purified form of the protein is injected. For instance, the PMPE polypeptide is resolved from other proteins using SDS-PAGE according to standard techniques well known to those skilled in the art, as previously described (Laemmli, 1970, Nature 227:680-685), and cutting the PMPE-containing band out of the gel. The excised band containing PMPE is macerated and injected into an animal to generate antiserum to the PMPE polypeptide. The antisera is examined using well known and generally accepted methods of ELISA to determine titer, Western blots to

determine binding to proteins, and for immunofluorescent staining and for complement-mediated cytotoxic activity against *Chlamydia*.

To aid in the characterization of the PMPE or PMPI protein, hyperimmune rabbit antisera is raised against whole EBs from *C. trachomatis*. Each animal is given a total of three immunizations of about 250 μg *Chlamydia* EBs per injection (beginning with the EBs mixed with complete Freund's adjuvant and followed with EBs mixed with incomplete Freund's adjuvant) at approximately 21 day intervals. At each immunization, approximately half of the material is administered intramuscularly (i.m.) and half is injected intranodally. Fourteen days after the third vaccination, a fourth booster of about 100 μg of EBs is given i.m. and the animals exsanguinated 7-10 days later.

6.5 ELISA

Anti-PMPE or anti-PMPI antibody titers are measured by ELISA using purified PMPE or PMPI protein (~1 µg/well) or *C. trachomatis* EBs (whole or crude protein extracts) or cells infected with *Chlamydia* as capture ligands by any method known by those skilled in the art. Serial dilutions of antisera are made in PBS and tested by ELISA in duplicate. HRP-conjugated antibody is diluted and used as the second reporter antibody in these assays. Titers are expressed as the greatest dilution showing positive ELISA reaction, *i.e.*, an O.D. 450 value >2SD above the mean negative control value (pre-bleed rabbit sera).

6.6 WESTERN BLOTS

Chlamydia trachomatis is grown in McCoy cells and Chlamydia cell lysates are prepared as described in section 6.1, supra. The solubilized cells are resolved on 12%
25 polyacrylamide gels and the separated proteins were electrophoretically transferred to PVDF membranes at 100 V for 1.5 hours as previously described (Thebaine et al. 1979, Proc. Natl. Acad. Sci. USA 76:4350-4354). The PVDF membranes are then pretreated with 25 ml of Dulbecco's phosphate buffered saline containing 0.5% sodium casein, 0.5% bovine serum albumin and 1% goat serum. All subsequent incubations are carried out using this
30 pretreatment buffer.

PVDF membranes are incubated with 25 ml of a dilution of preimmune rabbit serum or serum from a rabbit immunized with PMPE or PMPI polypeptide (as described above) for 1 hour at room temperature or with monoclonal antibodies to PMPE or PMPI. PVDF membranes are then washed twice with wash buffer (20 mM Tris buffer [pH 35 7.5.] containing 150 mM sodium chloride and 0.05% Tween-20). PVDF membranes are

incubated with 25 ml peroxidase-labeled goat anti-rabbit (or anti-mouse for monoclonals) IgG (Jackson ImmunoResearch Laboratories, West Grove, PA.) for 30 minutes at room temperature. PVDF membranes are then washed 4 times with wash buffer, and are developed with 3,3'diaminobenzidine tetra-hydrochloride and urea peroxide as supplied by Sigma Chemical Co. (St. Louis, Mo.; catalog number D-4418) for 4 minutes each.

Hyperimmune antisera or monoclonal antibody is used to probe Western blots of crude EB or RB extracts as well as 1.0% OGP EB extract preparations to identify PMPE or PMPI polypeptides from other *C. trachomatis* serovars and *Chlamydia* species. Lysates from *C. trachomatis* A, B, Ba, C, D, Da, E, F, G, H, I, Ia, J, K, L1, L2, L3, or MoPn serovars or *Chlamydia pneumoniae* are electrophoresed to identify proteins reactive with antisera generated against PMPE or PMPI protein.

6.7 CELLULAR ENVELOPE LOCATION OF PMPE AND PMPI

Surface localization of the PMPE or PMPI protein on different *Chlamydia*15 strains and derivatives is examined by indirect fluorescence antibody (IFA). IFA is
performed using the procedures generally known by those skilled in the art using
hyperimmune anti-PMPE or PMPI protein as the primary antibody. Hak cells infected with
whole EBs from *C. trachomatis* are achieved by the following method.

McCoy or Hak cells are grown to confluence in D-MEM media on 12 mm
20 plain coverslips inside 24 well tissue culture plates then centrifugally inoculated with ~5 X
10⁴ inclusion forming units (IFU) of the *C. trachomatis*. After ~24 hours incubation, the
culture media is removed and infected cells fixed in methanol for 10 min. The fixed
monolayer is then washed with PBS (1X) to remove fixative and overlayered with >300 μl
of anti-PMPE or PMPI rabbit antibody that has been diluted in PBS. After 1 hour
25 incubation with the primary antibody, the cells are washed gently with PBS then incubated
for ~30 minutes with mouse anti-rabbit IgG antibody conjugated with FITC. The second
antibody is diluted using a PBS solution containing 0.0091% Evans Blue as a counter stain
to visualize the monolayer. Cells are washed 2X in PBS to remove the secondary antibody,
the coverslips are removed from the culture plates, and mounted onto microscope slides
30 using a fluorescent mounting medium.

Identical cell samples are stained with pre-bleed rabbit antibody or FITC-conjugated second antibody alone are processed in parallel and served as antibody specificity (negative) controls. Counterstained samples are examined at a 1000-X magnification with a Zeiss Axioskop photomicroscope equipped with plan-neoflur objectives.

6.8 PROPERTIES OF PMPE and PMPI POLYPEPTIDES

PMPE polypeptide exists as a protein of approximately 90-115 KDa in its native state as determined via Western blots of crude EB or RB extracts of *Chlamydia*, as described in Section 6.1. PMPI polypeptide exists as a protein of approximately 90-115 KDa in its native state as determined via Western blots of crude EB or RB extracts of *Chlamydia*.

The isoelectric point of the PMPE protein is about 7.17. The isoelectric point of the PMPI protein is about 6.36.

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6.9 VACCINE EFFICACY

An *in vitro* neutralization model using methods generally known to those skilled in the art is used to show that protective antiserum inhibits Chlamydial infection (neutralization) of various tissue culture cell lines. Animal models are also essential for testing vaccine efficacy with both small animal (non-primate) and primate models necessary for preclinical evaluation. The guinea-pig is used for studying experimental ocular and genital infection by the Guinea-pig inclusion conjunctivitis agent (GPIC), *C. psittaci*.

The mouse offers a consistent and reproducible model of genital tract infection using human genital tract isolates. This mouse model is a generally accepted pre20 clinical assay and was used to evaluate MOMP as a subunit vaccine. Another model is known as the primate model of trachoma infection wherein the induction of secretory IgA was shown to be a prime component of protection. Vaccinogenic ability of new subunit antigen candidates is determined using the above-mentioned generally accepted *in vitro* neutralization and animal model systems.

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6.9.1 IN VITRO NEUTRALIZATION MODEL

As a preliminary exercise to the animal protection studies, hyperimmune anti-PMPE or PMPI antibody is evaluated for its ability to block the infectivity of various *C. trachomatis* serovars (*e.g.*, L2, B, F) *in vitro*. Although McCoy cells are used to propagate 30 *Chlamydia*, these cells also allow antibody-mediated uptake via Fc receptors. Therefore, to evaluate anti-PMPE or anti-PMPI antibody inhibition of infectivity, Hak cells, which do not display Fc receptors, are used in these analyses.

Cells are grown on coverslips in 24-well plates to a subconfluent monolayer (about 90% confluency = 1 X 10⁵ cells/ml) at 37°C in 5% CO₂. Anti-PMPE or PMPI antibody is diluted to about 100 µg/ml (total protein) in sucrose-phosphate-glutamate (SPG)

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buffer and then serially diluted in SPG buffer. Frozen aliquots of pretitered Chlamydia EBs are diluted in SPG buffer to about 2 X 104 IFU (inclusion forming units)/ml. EBs are premixed with the diluted anti-PMPE or PMPI antibody and incubated 30 minutes at 37°C on a rocking platform.

Prepared Hak cells are washed in HBSS and then incubated with the anti-PMPE or PMPI antibody/Chlamydia EB mixture in triplicate for each antibody using ~500 IFU/ml. Plates are centrifuged at ~500 rpm at room temperature for 1 hour then incubated for 2 hours at 37°C or are incubated for 2 hours at 37°C without prior centrifugation to allow EB infection of the monolayer. Then the inoculum is removed and plates washed 3 10 times with HBSS. Tissue culture media containing 1 μg/ml of cycloheximide is added and plates incubated for about 24-48 hours at 37°C in 5% CO₂ to allow inclusion bodies to develop. After incubation, the media is removed and cell monolayers washed 3X in PBS. Plates are fixed in methanol for 20 minutes and re-washed in PBS.

Cells are stained to visualize inclusions by incubating with anti-Chlamydia 15 LPS antibody (diluted about 1:500), washed 3 times in PBS, followed by incubation with FITC-conjugated goat secondary antibody for 30 minutes at 37°C. Coverslips are washed, air dried, and mounted in glycerol on glass slides. Inclusions are counted in five fields through the midline of the coverslip on a Zeiss fluorescence photomicroscope. Results are reported as the percent reduction of inclusion-containing cells with respect to a

20 heterogenous antibody control (rabbit pre-bleed sera).

6.9.2 VACCINE EFFICACY (MOUSE MODEL OF SALPINGITIS AND **FERTILITY**)

IMMUNIZATION AND CHALLENGE

25 The Tuffrey murine infertility model is employed to evaluate the efficacy of rPMPE or rPMPI to protect animals against Chlamydia trachomatis-induced salpingitis and infertility. Three groups of female C3HeOuJ mice (6 weeks of age, Jackson Labs) are employed for this evaluation. The test group is immunized (e.g., intradermally, intraperitoneal, subcutaneously, intramuscularly, or mucosally) by administration of a 30 vaccine formulation containing approximately 10-100 μg of purified rPMPE or rPMPI and adjuvant. For instance, mice that are immunized intranasally are sedated using an anesthesia cocktail consisting of 16% Ketaject and 16% Xlaject in 68% pyrogen-free PBS (100 µl) i.p./animal. Sedated animals are placed on their backs and using a standard laboratory pipette administered the vaccine formulations; approximately 10 µl of the 35 vaccine solution per nostril with a 5-10 minute wait period between applications.

Two groups of female mice are immunized similarly but with a preparation containing adjuvant only, no antigen. One of these groups is subsequently challenged with *C. trachomatis* (sham immunized, infected) and serves as the negative fertility control while the other group is not challenged (sham immunized, sham infected) and serves as the positive fertility control.

At week 4, all animals are administered a single i.p. dose of progesterone (2.5 mg in pyrogen-free PBS, Depo-Provera, Upjohn) to stabilize the uterine epithelium. At week 5, animals immunized with rPMPE or rPMPI and animals in the negative control group are infected by bilateral intrauterine inoculation with approximately 5 X 10⁵ IFU of C. 10 trachomatis (including but not limited to, from serovar F, strain NI1) in 100 μl of sucrose phosphate glutamate buffer (SPG). To mimic the manipulations to the reproductive tract experienced by the other groups, animals in the positive control are bilaterally inoculated with 100 µl of a McCoy cell extract that contains no C. trachomatis. At week 7, animals from each group are sacrificed by CO2 asphyxiation and the complete genital tract (both 15 upper and lower reproductive tracts) are removed for histopathological analysis. At week 9, the remaining females from each group are caged with 8-10 week old male C3H mice for a 2 month breeding period to assess fertility (1 male for every 2 females per cage with weekly rotation of the males within each group, animals from different experimental groups were not mixed). Palpation and periodic weighing are used to determine when animals in each 20 pair become pregnant. The parameters used to estimate group fertility are: F, the number of mice which littered at least once during the mating period divided by the total number of mice in that study group; M, the number of newborn mice (born dead or alive) divided by the number of litters produced in that group during the mating period; and N, the number of newborn mice (born dead or alive) divided by the total number of mice in that group.

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HISTOPATHOLOGY

Genital tracts are treated for > 24 hours in Bouin's fixative, progressively dehydrated in 50%, 70% and 100% methanol, soaked in toluol, and either paraffin embedded or directly embedded in OCT compound (Tissue-TEK, Miles) and subsequently snap frozen in liquid nitrogen. Tissue sections (approximately 6 µm) are stained with hematoxylin and eosin (after deparaffinization of the Bouin fixed samples). Inflammatory changes in the oviducts and ovaries are graded as follows: 0, no apparent inflammatory reaction; 1, a few mononuclear cells infiltrating the periovarial space or the submucosa of the oviduct; 2, same as 1 but to a greater extent; 3, same as 2 but with a thickened oviductal submucosa and the presence of inflammatory cells in the oviductal lumen; 4, same as 3 but

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to a greater extent. Inflammation in the cervix/vagina is scored based on the level of the intraepithelial infiltrate observed.

DETERMINATION OF rPMPE or rPMPI SPECIFIC HUMORAL RESPONSES

Blood samples are collected periodically during the immunization and challenge phases by retroorbital bleeding and serum prepared by centrifugation. Vaginal secretions are collected by repeated injection of 50-100 μ l of sterile PBS into the vagina with a standard laboratory pipetter and immediately withdrawing the solution. Two to three injection/withdrawal cycles are performed.

Ouantitation of antibody (Ab) responses by ELISA are performed as described in Section 6.5. Microwell ELISA plates (Maxisorb, NUNC) for determining antibody levels are coated overnight at 4°C with ~0.5-1.0 µg of purified rPMPE or rPMPI per well in 10 mM carbonate/bicarbonate buffer (pH 9.6), washed with PBS containing 0.1% Tween-20 (washing buffer) and blocked for ~1 hr at 37°C with a PBS solution containing 3% BSA. For the determination of antigen-specific serum IgG levels, test sera or vaginal secretions are serially diluted in washing buffer containing 0.5% BSA and aliquots (100 µl) incubated in the antigen-coated wells for ~2 hr at 37°C. The plates are then washed and incubated for ~1 hr at 37°C with a horseradish peroxidase (HRP)-conjugated goat antimouse IgG second antibody (Sigma). A HRP-conjugated goat anti-mouse IgA secondary antibody is used to detect the presence of rPMPE or rPMPI specific IgA in serum or vaginal secretions. After incubation with the appropriate secondary Ab, the plates are washed and incubated for ~20-30 minutes at room temperature with TMB substrate (Sigma). Reactions are stopped by the addition of 2M H₂SO₄ and the absorbance determined at 450 nm on a Molecular Devices SpectroMax microplate reader. Titers are determined as the reciprocal of the sample dilution corresponding to an optical density of 1.0 at 450 nm.

DETERMINATION OF rPMPE or rPMPI SPECIFIC CELLULAR RESPONSES

Groups of mice are immunized with rPMPE or rPMPI plus adjuvant vaccine as described above. At week 5, animals from each group are sacrificed by CO₂

asphyxiation, spleens removed and single cell suspensions prepared using conventional methodologies. For cellular assessment, animals are not treated with hormone. For both the positive control group (sham immunized and sham infected) and the negative control group (sham immunized, infected), spleen cells are pooled and tested for restimulation.

For the measurement of spleen cell proliferation, spleens are ground (5 to 10 rounds) in 5 ml RPMI 1640 Glutamax I supplemented with 10% fetal calf serum, 25 mM

HEPES, 50 U/ml penicillin, 50 μg/ml streptomycin, 1 mM sodium pyruvate, nonessential amino acids, and 50 mM 2-mercaptoethanol (Gibco-BRL). Live cells are counted by Trypan Blue staining and diluted in the same media to reach a density of 1.0 - 2.0 X 10⁶ cells/ml (Falcon 2063 polypropylene tubes). Triplicate cultures are set-up in round bottom 96-well culture plates (Nunclon, Nunc) using ~5 X 10⁵ responder cells per well in 200 μl of media. Cells are stimulated with rPMPE or rPMPI (antigen-specific proliferation) or with concanavalin A (Boerhinger Mannheim) as a positive stimulation control; unstimulated cell cultures are used as a negative control of cellular activation. After 72-96 hours of incubation at 37°C in 5% CO₂, cells are pulsed labeled for ~18 hrs with 1.0 Ci ³H-

10 thymidine (Amersham) per well. Pulsed cells are harvested onto glass-fiber sheets using a Tomtec Cell Harvester (Mk III) and counted for beta-emission in a 3-channel Wallac 1450 Trilux Liquid Scintillation Counter. The stimulation index (SI) for a sample (individual or pooled) is defined as the mean of the antigen or ConA-stimulated T-cell uptake of ³H-thymidine for triplicate wells divided by the mean of the unstimulated uptake for triplicate wells. SIs for both antigen-specific (rPMPE or rPMPI-specific) and ConA-specific proliferation are determined.

Results presented in Figure 7 demonstrate that animals immunized with recombinant PMPE protein develop a strong and uniform antigen-specific T-cell proliferative response. These data also show that T-cells from animals immunized with recombinant PMPE recognize and are strongly stimulated by the infectious EB form of *C. trachomatis*.

6.10 ISOLATION OF C. TRACHOMATIS CHROMOSOMAL DNA

Chlamydia trachomatis were grown as described supra in Section 6.1. Gradient purified EBs were suspended in ~5.0 ml of sterile water. An equal volume of lysis buffer (200 mM NaCl, 20 mM EDTA, 40 mM Tris-HCl pH 8.0, 0.5% (w/v) SDS, 0.5% (v/v) 2-mercaptoethanol, and 250 μg/ml of proteinase K) was added and the cells suspended by gentle agitation and trituration. The cell suspension was then incubated ~12 hours at 50°C to lyse the EBs and liberate chromosomal DNA. Proteinaceous material was precipitated by the addition of 5.0 ml of saturated NaCl (~6.0 M, in sterile water) and centrifugation at ~5,500 X g in a Sorval SS34 rotor at room temperature. Chromosomal DNA was precipitated from the cleared supernatant by the addition of two volumes of 100% ethanol. Aggregated DNA was collected and washed using gentle agitation in a small volume of a 70% ethanol solution. Purified chromosomal DNA was suspended in sterile water and allowed to dissolve/disburse overnight at 4°C by gentle rocking. The concentration of

dissolved DNA was determined spectrophotometrically at 260nm using an extinction coefficient of 1.0 O.D. unit ~50 µg/ml.

6.11 PCR CLONING OF THE PMPE ORF

- To produce high levels of recombinant PMPE protein for immunogenicity and protective efficacy studies, the PMPE ORF was PCR cloned into an *E. coli* high expression vector, pQE-30 (QiaGen). Genes cloned into the pQE-30 vector are expressed from a T5 promoter which is under the control of the lac operator. Genes cloned into pQE-30 are expressed as a fusion protein containing a Met-Arg-Gly-Ser-(His)₆ (SEQ ID NO.:71)
- 10 N terminus. An additional 15 amino acid segment encoded by vector sequences residing between the (His)₆ domain and the SalI site are also part of the N terminus. Oligonucleotide PCR primers complementary to the DNA sequences encoding the first 10 amino acid residues of the N-terminus of the mature protein and the last 9 C-terminal amino acid residues of the *C. trachomatis* serovar L2 PMPE ORF present in the UC Berkeley genomic database (http://chlamydia-www.berkeley.edu) were synthesized. In addition to the PMPE specific sequences, these PCR primers were designed to contain flanking SalI restriction
 - specific sequences, these PCR primers were designed to contain flanking SalI restriction sites to allow cloning of the ORF into the pQE-30 expression vector. The oligonucleotides (SEQ ID NOs.:66 and 67) were used to amplify PMPE-specific PCR products from *C. trachomatis* L2 serovar, genomic DNA.
- The sequence of the amplification primers for SEQ ID NO.:66 (forward primer) and SEQ ID NO.:67 (reverse primer) used for these PCR reactions are shown below.

Forward primer for PMPE

25 5'- ATC CAG CAG AG GGT CGA CGG GTT CCA GAT CCT ACG AAA GAG TCG CTA TC- 3' (SEQ ID NO.:66.)

Reverse primer for PMPE

5'- ATC CAG CAG AGG GTC GAC GGC C TTA GAA TCG CAG AGC AAT TTC CCC 30 ATT GA - 3' (SEQ ID NO.:67)

In addition to the PMPE coding sequences, a TAA (TAA in reverse complement) stop codon was engineered into the reverse primer immediately after the last PMPE codon to end protein synthesis. A SalI restriction site was engineered into the

reverse primer downstream and adjacent to the TAA stop codon to facilitate cloning into pQE-30.

Standard PCR amplification reactions (2 mM Mg²⁺, 200 μmol dNTPs, 0.75 units AmpliTaq, 50 μl final volume) were programmed using ~0.1 μg of *C. trachomatis* L2chromosomal DNA. Amplification of the PMPE target sequence was achieved using a standard 32-cycle, three-step thermal profile, *i.e.*, 95°C, 30 sec; 55°C, 45 sec; 72°C, 3 min. Amplification was carried out in 0.2 ml polypropylene thin-walled PCR tubes (Perkin-Elmer) in a Perkin-Elmer model 2400 thermal cycler. PCR amplification reactions produced the expected PMPE-specific ~3.0 Kbp amplimer.

10 The ~3.0 Kbp PMPE amplimer was purified from unincorporated primers using hydroxyapatite spin columns (QiaGen) and digested to completion with an excess of SalI (BRL, ~10 units per 1 μg DNA) according to the manufacturers recommendations. The purified and digested PMPE ORF was then purified *via* QiaGen columns as described above and cloned into plasmid pQE-30 that had been previously digested to completion with SalI and treated with calf intestinal alkaline phosphatase (BRL, ~0.05 units / pmole of 5' ends) to prevent vector religation (~5:1, insert:vector ratio).

Digestion of the pQE-30 vector with SalI produces a single restriction fragment of \sim 3.4 kbp.

Aliquots from the ligation reaction were then used to electrotransform a suitable *E. coli* host (*e.g.*, M15 (pREP)). Transformed cells were plated on 2X-YT agar containing 100 μg/ml ampicillin (for pQE-30) and 40 μg/ml kanamycin (for pREP) grown at 37°C for ~12-16 hours. Mini-prep DNA from ampicillin/kanamycin-resistant transformants picked at random were prepared using commercially available reagents (QiaGen Mini Prep Kit) and examined for the presence of recombinant plasmids larger than the pQE-30 plasmid (*i.e.*, insert-carrying plasmids). Putative PMPE-insert carrying recombinant plasmids were then digested to completion with SalI and examined for the presence of the PMP-specific fragment by standard agarose gel electrophoresis (0.8% agarose, TAE buffer). All plasmids tested were found to contain the PMPE insert. Plasmid M15 pREP (pQE-pmpE-CT)#37 was one recombinant derivative isolated by these

6.12 PCR CLONING OF THE PMPI ORF

To produce high levels of recombinant PMPI protein for immunogenicity and protective efficacy studies, the PMPI ORF was PCR cloned into an *E. coli* high expression vector pBAD/Thio-E (Invitrogen) under the control of the *ara*BAD promoter.

Oligonucleotide PCR primers complementary to the DNA sequences encoding the first 10 amino acid residues of the N-terminus and the last 8 C-terminal amino acid residues of the *C. trachomatis* L2 PMPI ORF present in the UC Berkeley genomic database (http://chlamydia-www.berkeley.edu) were synthesized. In addition to the PMPI specific sequences, the 5' forward PCR primer contained the sequence 5' AGG CAG AGG CAT-3' (SEQ ID NO.:68) immediately upstream of the PMPI ATG start codon while the 3' reverse primer contained the sequence 5'-AGG CAG AGG GTC GAC'3' (SEQ ID NO.:70) immediately downstream of the C-terminal GAA codon. The oligonucleotides (SEQ ID NO.:69 and 70) were used to amplify PMPI-specific PCR products from *C. trachomatis* L2 genomic DNA. The sequence of the amplification primers for SEQ ID NO.:69 (forward primer) and SEQ ID NO.:70 (reverse primer) used for these PCR reactions are shown below.

Forward primer PMPI

15 5'- AGG CAG AGG CAT ATG CGA CCT GAT CAC ATG AAC TTC TGT TG- 3' (SEQ ID NO.:69)

Reverse primer PMPI

5'-AGG CAG AGG GTC GAC GAA CCT GTA AGT GGT CCC CAG ATC -3' (SEQ ID NO.:70)

Standard PCR amplification reactions (2 mM Mg²⁺, 200 μmol dNTPs, 0.75 units AmpliTaq, 50 μl final volume) were programmed using ~0.1μg of *C. trachomatis* L2 chromosomal DNA. Amplification of the PMPI target sequence was achieved using a standard 32-cycle, three-step thermal profile, *i.e.*, 95°C, 30 sec.; 55°C, 45 sec.; 72°C, 3 min. Amplification was carried out in 0.2ml polypropylene thin-walled PCR tubes (Perkin-Elmer) in a Perkin-Elmer model 2400 thermal cycler. PCR amplification reactions produced the expected PMPI-specific ~3.0 Kbp amplimer.

The ~3.0 Kbp PMPI amplimer was purified from unincorporated primers using hydroxyapatite spin columns (QiaGen). The purified PMPI ORF was then cloned into the T/A PCR product cloning vector pUni/V5-HIS-TOPO plasmid (Invitrogen) using standard TOPO cloning methods described by the manufactor. The plasmid pUni/V5-His-TOPO carries a loxP site upstream of the T/A cloning site that can be used to recombine insert carrying plasmids with other loxP-containing plasmids for subsequent controlled

35 expression of recombinant protein.

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Aliquots from the PMPI amplimer and pUni/V5-His-TOPO ligation reaction were then used to transform a suitable E. coli host (e.g., PIR1). Transformed cells were plated on 2X-YT agar containing 40 μg/ml kanamycin and grown at 37°C for ~12-16 hours. Mini-prep DNA from kanamycin-resistant transformants picked at random were prepared using commercially available reagents (QiaGen Mini Prep Kit) and examined for the presence of recombinant plasmids larger than the ~2.2 Kbp vector plasmid Uni/V5-His -TOPO (i.e., insert-carrying plasmids). pUni/V5-His-TOPO plasmids carrying the cloned PMPI ORF were mixed (~100 ng) with the plasmid pBAD/Thio-E (~100 ng) in the presence of a recombination buffer supplied by the manufacturer and the two were recombined into a 10 single molecule by homologous recombination at the lox sites on both the pUni/V5-His-TOPO-PMPI plasmid and pBAD/Thio-E vector catalyzed by the Cre recombinase. pBAD/Thio-E carries a segment of the E. coli thioredoxin protein (~12 Kda) that has been mutated to encode an N-terminal (His) affinity chromatography domain. A single loxP recombination site is also encoded by pBAD/Thio-E immediately downstream of the His-15 Thio redoxin coding sequence. The His-Thioredoxin coding sequence and the loxP site are themselves downstream of the araBAD promoter. Recombining pBAD/Thio-E with PMPI derivatives of pUni/V5-His-TOPO through the loxP sites using the loxP-specific Cre recombinase results in the formation of a chimeric protein with the His-thioredoxin domain at the N-terminus which is fused in frame to the PMPI ORF. Following recombination the 20 chimeric plasmids were transformed into E. coli TOP10 cells. Transformed cells were selected on plates containing 40 µg/ml kanamycin. All ~9.5 kBp plasmids tested were found to contain the PMPI insert. Plasmid TOP10 (pBAD-PmpI-Ct-Uni)#7 was one recombinant derivative isolated by these procedures.

25 6.13 EXPRESSION OF RECOMBINANT PMPE

One milliliter of a frozen stock of *E. coli* strain M15 pREP containing plasmid (pQE-pmpE-CT)#37 was used to inoculate ~100 ml of 2X-YT broth containing 40 µg/ml kanamycin and 100 µg/ml ampicillin and grown overnight at 30°C to prepare a fermentor seed culture. Approximately 20 ml of the overnight seed culture was then used to inoculate a New Brunswick Bioflow 3000 fermentor loaded with ~2.0 L of 2X-YT broth containing 40 µg/ml kanamycin and 100 µg/ml ampicillin. The culture was grown at 37°C with vigorous aeration until an O.D.600 value of ~1.0 was attained. Expression of rPMPE was induced by adding IPTG to a 1.0 mM final concentration and continuing fermentation. Incubation in the presence of ITPG was continued for approximately 4-5 hours.

At the end of the induction period, the E. coli culture, with some cells displaying classic recombinant protein inclusion bodies, was harvested by continuous flow centrifugation using an Heraeus Contifuge 28RS centrifuge. Following centrifugation, cell mass was scraped from the centrifuge bowl and stored at -70°C until processed.

Approximately 15 gm of the M15 pREP (pQE-pmpE-CT)#37 frozen cell paste was resuspended by vortexing and trituration in ~40 ml of ice cold 10 mM sodium phosphate buffer, pH 7.3. Once suspended, lysozyme (Chicken egg white, Sigma) and DNase I (Bovine pancreas, Sigma) were added to final concentrations of 1.0 mg/ml and 0.01 mg/ml, respectively, and the mixture incubated on ice for 30-45 minutes. Cells were 10 disrupted by 2 sequential passes through a pre-cooled (~4°C) SLM Aminco French Pressure Cell (~14 Kpsi, 1" diameter bore). The cell lysate was then centrifuged for 5 min at ~500 X g (4°C) in a Sorvall SS34 rotor to remove unbroken cells. Insoluble material containing the rPMPE was isolated (pelleted) by centrifugation for 45 minutes at ~20,000 X g (4°C) in a Sorvall SS34 rotor. The supernatant from this centrifugation was discarded and the 15 insoluble fraction stored at -20°C in pellet form.

To selectively extract contaminating proteins and remove endotoxin, the rPMPE-containing insoluble pellet was thawed on ice and washed twice with 10 ml of PBS buffer containing 2.0% Triton X-100. Washing was performed at room temperature and suspension of the gelatinous rPMPE-containing pellet was accomplished by vortexing and 20 homogenization in a conventional glass tissue grinder. Insoluble material containing the rPMPE was recovered after washing by centrifugation at ~10,000 X g for 20 minutes (room temperature) in a Sorvall SS34 rotor. Insoluble material was then washed (again by vortexing and homogenization) 2 times with 10 ml of a 4.0 M urea solution containing 2.0 M NaCl. Washed rPMPE material was recovered by centrifugation as above. The insoluble 25 rPMPE fraction was further washed 2 times with 10 ml of a PBS solution containing 1.0% Zwittergent 3-14 (Sigma).

The rPMPE pellet recovered after centrifugation of the final wash solution was then solubilized for 2 hours at room temperature in standard Laemelli SDS-PAGE sample buffer containing 4 M urea. Solubilized rPMPE was size fractionated into a single 30 protein band of ~ 105 kDa by electrophoresis through a cylinderical ~ 37 mm X ~ 12 cm 6% polyacrylamide (36:1, acrylamide:bis-acrylamide) Tris/glycine/SDS preparative gel. A 4% polyacrylamide stacking gel was formed on top of the resolving gel (~37mm X ~3cm). Electrophoresis was carried out on a BioRad model 491 PrepCell unit for ~12 hours at ~22°C (12 watts constant wattage) using a conventional Tris/glycine/SDS running buffer 35 (BioRad). As electrophoresis proceeded, size fractionated proteins eluting from the bottom

of the PrepCell were pumped to an Amersham fraction collector where ~8.0 ml fractions were collected.

Aliquots (~20µl) from each fraction were mixed with an equal volume of 2X LSB heated to ~100°C for 3 minutes and electrophoresed on a 4-20% SDS PAG to identify fractions containing the PMPE protein. Prestained molecular weight standards were loaded into a parallel lane and were used as a size gauge. PMPE-containing factors were pooled and excess SDS detergent was removed by diluting the eluted sample with approximately 20 volumes of sterile, endotoxin-free 10 mM sodium phosphate buffer (pH 7.4) and concentrated to approximately 1.0 mg/ml by ultrafiltration in an Amicon stirred concentration cell using a YM30 ultrafiltration membrane

Residual endotoxin was removed from the concentrated rPMPE solution by polymyxin B Affi-Prep Polymyxin Matrix (BioRad) treatment. Affi-Prep treatment was performed overnight at 4°C in a batch mode according to the manufacturers recommendations.

The protein concentration of the concentrated, polymyxin B-treated rPMPE was determined using the Micro BCA method (Pierce Chem.) and BSA as a standard.

Purified rPMPE (~0.9-1.2 mg/ml protein concentration) was evaluated for purity, identity, and residual endotoxin burden by SDS-PAGE, Western blot, and a colorimetric endotoxin assay (BioWhittaker), respectively. The gel-purified rPMPE 20 material displayed a purity of >95% as a single band of the expected molecular size by gel analysis. Residual endotoxin is calculated to be ≤ 0.05 EU/g.

6.14 EXPRESSION OF RECOMBINANT PMPI

One milliliter of a frozen stock of *E. coli* strain TOP10 containing plasmid

(pBAD-pmpI-Ct-Uni)#7 was used to inoculate ~100 ml of 2X-YT broth containing 40 µg/ml kanamycin and grown overnight at 37°C to prepare a fermenter seed culture.

Approximately 20 ml of the overnight seed culture was then used to inoculate a New Brunswick Bioflow 3000 fermenter loaded with ~2.0 L of 2X-YT broth containing 40 µg/ml kanamycin. The culture was grown at 37°C with vigorous aeration until an O.D.600

value of ~1.0 was attained. Expression of rPMPI was induced by addition of arabinose to a final concentration of 1.0 mM and continuing cultivation for approximately 4-5 hours.

At the end of the induction period, the *E. coli* culture, with some cells displaying classic recombinant protein inclusion bodies, was harvested by continuous flow centrifugation using an Heraeus Contifuge 28RS centrifuge. Following centrifugation, cell mass was scraped from the centrifuge bowl and stored at -70°C until processed.

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Approximately 15 gm of the TOP10 (pBAD-pmpI-Ct-Uni)#7 frozen cell paste was resuspended by vortexing and trituration in ~40 ml of ice cold 10mM sodium phosphate buffer, pH 7.3. Once suspended, lysozyme (Chicken egg white, Sigma) and DNAase I (Bovine pancreas, Sigma) were added to final concentrations of 1.0 mg/ml and 0.01 mg/ml, respectively, and the mixture incubated on ice for 30- 45 minutes. Cells were disrupted by 2 sequential passes through a pre-cooled (~4°C) SLM Aminco French Pressure Cell (~14 Kpsi, 1" diameter bore). The cell lysate was then centrifuged for 5 min at ~500 X g (4°C) in a Sorvall SS34 rotor to remove unbroken cells. Insoluble material containing the rPMPI was isolated (pelleted) by centrifugation for 45 min at ~20,000 X g (4°C) in a Sorvall SS34 rotor. The supernatant from this centrifugation was discarded and the insoluble fraction stored at -20°C in pellet form.

To selectively extract contaminating proteins and remove endotoxin, the rPMPI-containing insoluble pellet was thawed on ice and washed twice with 10 ml of PBS buffer containing 2.0% Triton X-100. Washing was performed at room temperature and suspension of the gelatinous rPMPI-containing pellet was accomplished by vortexing and homogenization in a conventional glass tissue grinder. Insoluble material containing the rPMPI was recovered after washing by centrifugation at ~10,000 X g for 20 minutes (room temperature) in a Sorvall SS34 rotor. Insoluble material was then washed (again by vortexing and homogenization) 2 times with 10 ml of a 4.0 M urea solution containing 2.0 M NaCl. Washed rPMPI material was recovered by centrifugation as above. The insoluble rPMPI fraction was further washed 2 times with 10 ml of a PBS solution containing 1.0% Zwittergent 3-14 (Sigma).

The rPMPI pellet recovered after centrifugation of the final wash solution was then solubilized for 2 hours at room temperature in standard Laemelli SDS-PAGE sample buffer containing 4 M urea. Solubilized rPMPI-thiroredoxin-fusion protein was size fractionated into a single protein band of ~ 105 kDa using a model 491 PrepCell (BioRad) cylinderal gel as described above for PMPE above. PMPI was obtained from SDS PAG and residual endotoxin removed as described for PMPE above.

The protein concentration of the concentrated, polymyxin B-treated rPMPI-30 thioredoxin fusion protein was determined using the Micro BCA method (Pierce Chem.) and BSA as a standard.

Purified rPMPI (~0.9-1.2 mg/ml protein concentration) was evaluated for purity, identity, and residual endotoxin burden by SDS-PAGE, Western blot, and a colorimetric endotoxin assay (BioWhittaker), respectively. The gel-purified rPMPI material

displayed a purity of >95% as a single band of the expected molecular size by gel analysis. Residual endotoxin is calculated to be ≤ 0.05 EU/g.

6.15 AFFINITY CHROMATOGRAPHY PURIFICATION OF RECOMBINANT PROTEIN

Recombinant PMPE or PMPI protein is purified to apparent homogeneity using standard preparative immobilized metal affinity chromatography (IMAC) procedures. Briefly, an *E. coli* strain harboring an expression plasmid containing PMPE or PMPI protein gene is grown in Luria broth in a 5L fermenter (New Brunswick) at 37°C with moderate aeration until mid-log phase (~0.5 O.D.₆₀₀) and induced with IPTG or arabinose (1 mM final) for 4-5 hours. Cell paste is collected, washed in PBS and stored at -20°C. Aliquots of frozen cell paste (~9-10 grams wet weight) are suspended in ~120 ml of D-PBS by mechanical agitation and lysed by passage through a French pressure cell (2X, 14,000 psi, 4°C). Soluble protein is then removed from rPMPE or rPMPI protein inclusion bodies by high speed centrifugation (~20,000 X g, 4°C, 30 min).

The insoluble pellet containing rPMPE or rPMPI protein is suspended in ~20 ml of ice cold D-PBS by homogenization and centrifuged as above. Washed rPMPE or rPMPI protein inclusion bodies are then denatured by suspension in a sodium phosphate buffer (0.1 M, pH 7.0) containing 7.4 M guanidine hydrochloride, 5% 2-ME and 10% glycerol and loaded onto a Ni²⁺-affinity column (1.5 cm X 25 cm, bed volume ~30 ml) prepared from Fast-Flow Chelating Sepharose (Pharmacia) and charged with Ni²⁺or Zn²⁺ ions by standard procedures. Unbound material is removed by washing the column with ~5-10 column volumes of a sodium phosphate buffer (0.1 M, pH 7.0) containing ~7.4 M guanidine HCl.

Recombinant PMPE or PMPI protein bound to the affinity resin by virtue of the (His)₆ affinity purification domain is eluted using sodium phosphate buffer (pH 7.4) containing 7.4 M guanidine HCl and 200 mM imidazole. Eluted material is dialyzed against TE buffer containing SDS (0.5%) to remove the guanidine. Dialyzed material is concentrated using a Amicon stirred cell concentrator using a YM30 membrane and mixed with a 1/5 volume of 5X SDS gel sample buffer containing 1 mM 2-mercaptoethanol and boiled at 100°C for 5 minutes.

Samples are loaded onto Tris/glycine acrylamide gels (4% stacking gel, 4-20% resolving gel, 30:0.8 acrylamide:bis solution, 1 mm thickness). A prestained molecular weight standard (SeeBlue or Multimark, Novex) is run in parallel with the rPMPE or rPMPI protein samples to identify size fractions on the gel. The purity of rPMPE

or rPMPI protein is determined using conventional SDS-PAGE and commercially available silver staining reagents (Silver Stain Plus, Novex).

6.16 GENERATION OF A RADIOLABELLED SCREENING PROBE

The sequence information shown above is used to design a pair of nondegenerate oligonucleotide primers. PCR amplification of DNA fragments is performed under the same conditions as described above with the exception that the annealing temperature is lowered to 50°C. The DNA fragment is isolated from an agarose gel as before and radiolabeled using [32P]-gamma-ATP and T4 polynucleotide kinase according to standard methods. Unincorporated radiolabel is separated from the probe on a G25 Sepharose spin column. Before use, the probe is denatured for 2 min. at 95°C and subsequently chilled on ice (4°C).

6.17 HYBRIDIZATION OF PLAQUE-LIFT FILTERS AND SOUTHERN BLOTS WITH RADIOLABELLED PROBE

Phage plaques from library platings are immobilized on nylon filters using standard transfer protocols well known to those skilled in the art. Digested bacterial genomic DNA, phage or plasmid DNA is electrophoresed on 0.8% TAE-agarose gels and transferred onto nylon filters using a pressure blotter (Stratagene) according to the manufacturer's recommendations. Hybridizations with selected probes are performed at 37°C (for example, to detect PMP homologs of other species). Hybridizations with specific probes are generally carried out at 50-60°C (e.g., to detect the identical sequence or the analogous protein in another serovan). Washes of increasing stringency are done at the respective hybridization temperatures until nonspecific background is minimized.

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6.18 CONSTRUCTION OF A Chlamydia GENOMIC DNA LIBRARY

A genomic library was constructed in the λZAPII replacement vector obtained from Stratgene. The vector arms were digested with EcoR1. Digests of *Chlamydia* DNA by EcoR1 were performed to yield fragment sizes between 1 kb and 5 kb.

30 Ligations of vector arms and insert DNA were carried out according to standard protocols. Ligation reactions were packaged *in vitro* using the Stratagene GigaPack Gold III extract. The packaged phage were plated on *E. coli* Xl Blue MRA (P2) (Stratagene). An initial library titer was determined and expressed as number of pfu.

The library is screened using 4 x 10⁴ pfu that are plated at a density of 8 x 10³ pfu/130 mm plate with a PMPE or PMPI specific, probe. Several putative positive phage

plaques are located and the strongest hybridizing phage are eluted from cored agarose plugs, titered and replated for secondary screening. The selected phages are replated at low density (approximately 100 pfu/plate) and plaques are analyzed by PCR using primer pairs. Inserts carrying plasmids (phagemids) are rescued from the selected phage by co-infecting *E. coli* cells with an appropriate helper virus.

6.19 <u>DETERMINATION OF INSERT SIZE AND MAPPING OF DNA</u> FRAGMENTS

In order to estimate the size of inserts, phagemid DNA is digested with appropriate restriction enzymes (e.g., NotI) and the digests are analyzed on a 0.5% TAE-agarose gel side by side with suitable DNA markers. In order to map restriction fragments that would hybridize to the probe, DNA from phagemid isolates is digested with a number of common restriction enzymes either alone or in combination with NotI. The rationale of this approach is to discriminate between fragments that span the insert/phagemid vector junction and those that map on the NotI insert. The series of single and double digests are run side-by-side for each phage isolate and analyzed by Southern analysis with radiolabeled probe.

6.20 SEQUENCING OF THE PMPE and PMPI GENE

Sequencing of the PMPE and PMPI gene is performed using the Dye Terminator Cycle Sequencing Kit from Perkin-Elmer according to the manufacturer's specifications. The sequencing reactions are read using an ABI Prism 310 Genetic Analyzer. The sequences are aligned using the AutoAssembler software (Perkin-Elmer) provided with the ABI Prism 310 sequencer.

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The present invention is not to be limited in scope by the microorganism deposited or the specific embodiments described herein. It will be understood that variations which are functionally equivalent are within the scope of this invention. Indeed, various modifications of the invention, in addition to those shown and described herein, will become apparent to those skilled in the art from the foregoing description and accompanying drawings. Such modifications are intended to fall within the scope of the appended claims.

Various publications are cited herein, the disclosures of which are incorporated by reference in their entireties.

WHAT IS CLAIMED IS:

- An isolated PMPE or PMPI polypeptide of a *Chlamydia spp.*, having a molecular weight between 90 and 115 kDa as determined by SDS polyacrylamide gel
 electrophoresis.
 - 2. The PMPE or PMPI polypeptide of claim 1, wherein the *Chlamydia* spp. is *Chlamydia trachomatis*, *Chlamydia pneumonia*, *Chlamydia psittaci* or *Chlamydia pecorum*.

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- 3. The PMPE or PMPI polypeptide of claim 2, wherein the *Chlamydia spp.* is *C. trachomatis*.
- 4. The PMPE or PMPI polypeptide of claim 1, which comprises an amino acid sequence of SEQ ID NO.:2 or 4, a sequence substantially homologous thereto, or an at least 8 amino acid fragment thereof.
- 5. The PMPE or PMPI polypeptide of claim 1, or a peptide fragment thereof, which specifically binds an antibody that specifically binds to a protein having the 20 amino acid sequence of SEQ ID NO.:2 or 4.
 - 6. A peptide fragment of the PMPE or PMPI polypeptide of claim 1, which fragment is at least 8 amino acids in length.
- 7. The peptide fragment of claim 6 wherein said peptide fragment comprises the amino acid sequence of one of SEQ ID NO.:5-34.
- 8. An isolated fusion polypeptide comprising at least two peptides, said at least two peptides each consisting of an amino acid sequence of one of SEQ ID NO.:5, 6, 30 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17,18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, or 34, with the proviso that the peptides are arranged in a configuration that is different from the configuration of a naturally occurring PMPE or PMPI polypeptide.

- 9. The isolated fusion polypeptide of claim 8 wherein the fusion polypeptide comprises peptides consisting of the amino acid sequences of SEQ ID NO.:5, 6, 7, 8, 9, 10 and 11.
- 5 10. The isolated fusion polypeptide of claim 8 wherein the fusion polypeptide comprises peptides consisting of the amino acid sequences of SEQ ID NO.:23, 24, 25, 26, 27, 28, and 29.
- 11. An antibody or an antigen-binding fragment thereof that specifically binds the PMPE or PMPI polypeptide of claim 1.
 - 12. An antibody or an antigen-binding fragment thereof that specifically binds the peptide fragment of claim 6.
- 13. An antibody or an antigen-binding fragment thereof that specifically binds a peptide fragment consisting of an amino acid sequence of SEQ ID NO.:5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17,18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, or 34.
- 20 14. The antibody of claim 11, 12 or 13 which is a cytotoxic, cytostatic, or neutralizing antibody.
 - 15. A vaccine comprising the PMPE or PMPI polypeptide of claim 1 and a pharmaceutically acceptable carrier or diluent.
 - 16. The vaccine of claim 15 further comprising one or more adjuvants or immunostimulatory compounds.
- 17. The vaccine of claim 16 wherein the adjuvants or immunostimulatory30 compounds are one or more of alum, MLT, QS21, MF59, CpG DNA, PML, calcium phosphate and PLG.
 - 18. The vaccine of claim 16 comprising one adjuvant or immunostimulatory compound.

- 19. The vaccine of claim 16 comprising two different adjuvants or immunostimulatory compounds.
- 20. A vaccine comprising the polypeptide fragment of claim 6 and a pharmaceutically acceptable carrier or diluent.
 - 21. The vaccine of claim 20 further comprising one or more adjuvants or immunostimulatory compounds.
- 10 22. The vaccine of claim 21 wherein the one or more adjuvants or immunostimulatory compounds are selected from the group consisting of alum, MLT, QS21, MF59, CpG DNA, PML, calcium phosphate and PLG.
- 23. The vaccine of claim 21 comprising one adjuvant or 15 immunostimulatory compound.
 - 24. The vaccine of claim 21 comprising two different adjuvants or immunostimulatory compounds.
- 20 25. A vaccine comprising the isolated fusion polypeptide of claim 8 and a pharmaceutically acceptable carrier or diluent.
 - 26. The vaccine of claim 25 further comprising one or more adjuvants or immunostimulatory compounds.
 - 27. The vaccine of claim 26 wherein the one or more adjuvants or immunostimulatory compounds are selected from the group consisting of alum, MLT, QS21, MF59, CpG DNA, PML, calcium phosphate and PLG.
- 30 28. The vaccine of claim 26 comprising one adjuvant or immunostimulatory compound.
 - 29. The vaccine of claim 26 comprising two different adjuvants or immunostimulatory compounds.

- 30. A vaccine comprising the antibody of claim 11 and a pharmaceutically acceptable carrier or diluent.
- 31. The vaccine of any one of claims 15, 20 or 25 additionally comprising one or more immunogens selected from the group consisting of a lipid, lipoprotein, phospholipid, lipooligosaccharide, protein, attenuated organism and inactivated whole cell.
- 32. The vaccine of claim 31 wherein the one or more immunogens are a DPT vaccine, a HMWP of *C. trachomatis*, a MOMP of *C. trachomatis*, or an entire organism, or subunit therefrom, of *Chlamydia*, *Neisseria gonorrhea*, HIV, *Haemophilus influenzae*, *Moraxella catarrhalis*, *Human papilloma virus*, *Herpes simples virus*, *Haemophilus ducreyi*, *Treponema pallidium*, *Candida albicans* or *Streptococcus pneumoniae*.

33. An isolated nucleic acid molecule comprising a nucleotide sequence encoding a PMPE or PMPI polypeptide, or an at least 8 amino acid fragment thereof, of a *Chlamydia spp.*, said PMPE or PMPI polypeptide having a molecular weight between 90 and 115 kDa as determined by SDS polyacrylamide gel electrophoresis.

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- 34. An isolated nucleic acid molecule comprising the nucleotide sequence of SEQ ID NO.:1 or 3, an at least 15 nucleotide fragment thereof, or the complement thereof.
- 25 35. A pharmaceutical composition comprising the isolated nucleic acid molecule of claim 33.
 - 36. A vaccine comprising the isolated nucleic acid molecule of claim 33.
- 37. A vaccine comprising an isolated nucleic acid comprising a nucleotide sequence encoding the PMPE or PMPI polypeptide of claim 1, or an at least 15 nucleotide fragment thereof, and further comprising one or more adjuvants or immunostimulatory compounds.

- 38. The vaccine of claim 37 wherein the one or more adjuvants or immunostimulatory compounds are alum, MLT, QS21, MF59, CpG DNA, PML, calcium phosphate or PLG.
- 5 39. The vaccine of claim 37 comprising one adjuvant or immunostimulatory compound.
 - 40. The vaccine of claim 37 comprising two different adjuvants or immunostimulatory compounds.

- 41. A vaccine comprising one or more of an isolated PMPE or PMPI polypeptide of a *Chlamydia spp.*, having a molecular weight between 90 and 115 kDa as determined in SDS polyacrylamide gel electrophoresis; or an isolated nucleic acid comprising a nucleotide sequence encoding an PMPE or PMPI polypeptide of a *Chlamydia spp.*, said PMPE or PMPI polypeptide having a molecular weight between 90 and 115 kDa as determined by SDS polyacrylamide gel electrophoresis; said vaccine further comprising one or more adjuvants or immunostimulatory compounds selected from the group consisting of alum, MLT, QS21, MF59, CpG DNA, PML, calcium phosphate and PLG.
- 42. A method of producing an immune response in an animal comprising administering to the animal an immunogenic amount of the PMPE or PMPI polypeptide of claim 1.
- 43. A method of producing an immune response in an animal comprising administering to the animal an immunogenic amount of the peptide fragment of claim 6.
 - 44. A method of producing an immune response in an animal comprising administering to the animal an immunogenic amount of the isolated fusion polypeptide of claim 8.

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45. A method of producing an immune response in an animal comprising administering to the animal an immunogenic amount of the nucleic acid molecule of claims 33 or 34.

- 46. A method of producing an immune response in an animal comprising administering to the animal an immunogenic amount of the vaccine of claim 41.
- 47. A method of producing an immune response in an animal comprising administering to the animal an immunogenic amount of one or more of the vaccines of claims 15, 20, 25, 30, 36, or 37, wherein said vaccines are administered simultaneously or sequentially.
- 48. Plasmid M15 pREP (pQE-pmpE-Ct-Uni)#37 obtainable from *E. coli*, as deposited with the ATCC and assigned accession number PTA-2462.
 - 49. Plasmid TOP10 (pBAD-pmpI-Ct-Uni)#7 obtainable from *E. coli*, as deposited with the ATCC and assigned accession number PTA-2461.
- 15 50. A recombinant expression vector adapted for transformation of a host cell comprising the nucleic acid molecule of claim 33 or 34.
- 51. The recombinant expression vector of claim 50 further comprising an expression means operatively coupled to the nucleic acid molecule for expression by the host of the PMPE or PMPI protein, or a fragment thereof.
 - 52. The expression vector of claim 51, wherein the expression means includes a nucleotide sequence encoding an amino acid sequence that can be used for the purification of the PMPE or PMPI protein.
 - 53. The expression vector of claim 51 wherein the expression means further includes a nucleotide sequence encoding an amino acid sequence that directs secretion from the host of the PMPE or PMPI polypeptide.
- 30 54. A transformed host cell containing the expression vector of claim 50.
 - 55. A transformed host cell containing the plasmid of claim 48 or 49.
- 56. A host cell containing the nucleic acid molecule of claim 33 or 34 operatively linked to a heterologous promoter.

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57. An isolated recombinant PMPE or PMPI polypeptide produced by a method comprising culturing the transformed host cell of claim 54 under conditions suitable for expression of said PMPE or PMPI polypeptide and recovering said PMPE or PMPI polypeptide.

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58. An isolated recombinant PMPE or PMPI polypeptide produced by a method comprising culturing the transformed host of claim 55 under conditions suitable for expression of said PMPE or PMPI polypeptide and recovering said PMPE or PMPI polypeptide.

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- 59. An isolated PMPE or PMPI polypeptide produced by a method comprising culturing the host cell of claim 56 under conditions suitable for the expression of a PMPE or PMPI polypeptide and recovering said PMPE or PMPI polypeptide.
- 15 60. A method of preventing, treating or ameliorating a disorder or disease associated with infection of an animal with *Chlamydia* by administering to a subject in need of such prevention, treatment or amelioration an effective amount of the polypeptide of claim 1.
- 20 61. A method of preventing, treating or ameliorating a disorder or disease associated with infection of an animal with *Chlamydia* by administering to a subject in need of such prevention, treatment or amelioration an effective amount of the polypeptide fragment of claim 6.
- A method of preventing, treating or ameliorating a disorder or disease associated with infection of an animal with *Chlamydia* by administering to a subject in need of such prevention, treatment or amelioration an effective amount of the isolated fusion polypeptide of claim 8.
- 30 63. A method of preventing, treating or ameliorating a disorder or disease associated with infection of an animal with *Chlamydia* by administering to a subject in need of such prevention, treatment or amelioration an effective amount of the vaccine of claim 30.

64. A method of preventing, treating or ameliorating a disorder or disease associated with infection of an animal with *Chlamydia* by administering to a subject in need of such prevention, treatment or amelioration an effective amount of the vaccine of claim 31.

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65. A method of preventing, treating or ameliorating a disorder or disease associated with infection of an animal with *Chlamydia* by administering to a subject in need of such prevention, treatment or amelioration an effective amount of the vaccine of claim 32.

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66. A method of preventing, treating or ameliorating a disorder or disease associated with infection of an animal with *Chlamydia* by administering to a subject in need of such prevention, treatment or amelioration an effective amount of the vaccine of claim 37.

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67. A method of preventing, treating or ameliorating a disorder or disease associated with infection of an animal with *Chlamydia* by administering to a subject in need of such prevention, treatment or amelioration an effective amount of the vaccine of claim 41.

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- 68. A method of preventing, treating or ameliorating a disorder or disease associated with infection of an animal with *Chlamydia* by administering to a subject in need of such prevention, treatment or amelioration an effective amount more than one of the vaccines of claims 15, 20, 25, 30, 31, 35, 37 or 41, each optionally comprising one or more immunogens selected from the group consisting of a lipid, lipoprotein, phospholipid, lipooligosaccharide, protein, attenuated organism and inactivated whole cell, wherein said vaccines are administered simultaneously or sequentially
- 69. An antagonist which inhibits the activity of the polypeptide of 30 claim 1.
 - 70. An antagonist which inhibits the expression of the nucleic acid of claim 33.

- 71. A method for identifying compounds that interact with the polypeptide of claim 1, said method comprising contacting a composition comprising the polypeptide with the compound to be screened under conditions that permit interaction between the compound and the polypeptide; and detecting the interaction of the compound with the polypeptide.
- 72. A method for identifying compounds which interact with the nucleic acid molecule of claim 33, said method comprising contacting a composition comprising the nucleic acid molecule with the compound to be screened under conditions that permit interaction between the compound and the nucleic acid molecule; and detecting the interaction of the nucleic acid with the compound.

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ABSTRACT OF THE DISCLOSURE

The invention discloses the *Chlamydia* PMPE and PMPI polypeptide, polypeptides derived thereof (PMP-derived polypeptides), nucleotide sequences encoding said polypeptides, antibodies that specifically bind the PMP polypeptides and PMP-derived polypeptides and T-cells specific for PMP polypeptides and PMP-derived polypeptides. Also disclosed are prophylactic and therapeutic compositions, including immunogenic compositions, *e.g.*, vaccines, comprising PMP polypeptides or PMP-derived polypeptides or antibodies thereto. The invention additionally discloses methods of inducing in animals an immune response to *Chlamydia* cells, *Chlamydia* elementary bodies, and/or cells expressing *Chlamydial* proteins, *e.g.*, cell infected with *Chlamydia*.

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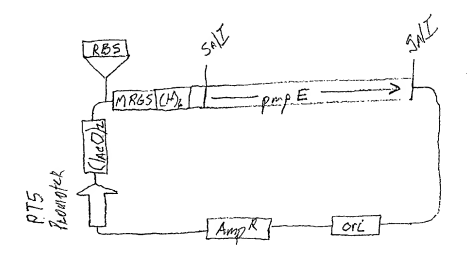


FIG. 1

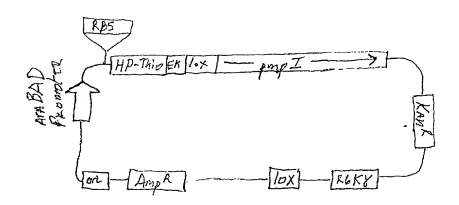


FIG. 2

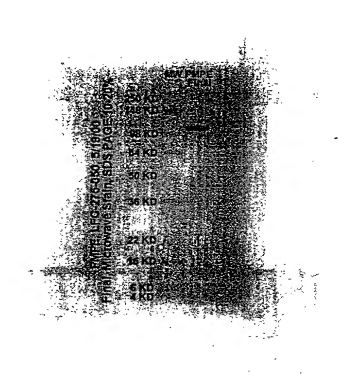


FIG. 3

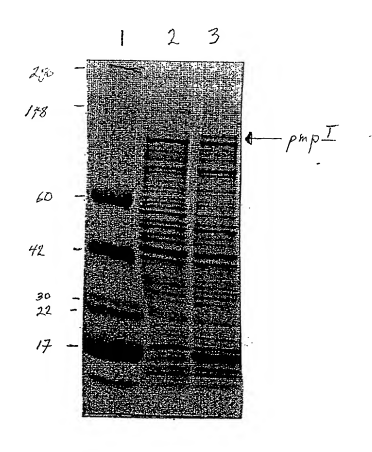


FIG. 4

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atg Met 1	aaa Lys	aaa Lys	gcg Ala	ttt Phe 5	ttc Phe	ttt Phe	ttc Phe	ctt Leu	att Ile 10	gga Gly	aac Asn	tcc Ser	cta Leu	tca Ser 15	gga Gly	48
											atg Met					96
cca Pro	gat Asp	cct Pro 35	acg Thr	aaa Lys	gag Glu	tcg Ser	cta Leu 40	tca Ser	aat Asn	aaa Lys	att Ile	agt Ser 45	ttg Leu	aca Thr	gga Gly	144
gac Asp	act Thr 50	cac His	aat Asn	ctc Leu	act Thr	aac Asn 55	tgc Cys	tat Tyr	ctc Leu	gat Asp	aac Asn 60	cta Leu	cgc Arg	tac Tyr	ata Ile	192
											gct Ala					240
aca Thr	gat Asp	tac Tyr	cta Leu	agc Ser 85	ttt Phe	ttt Phe	gat Asp	aca Thr	caa Gln 90	aaa Lys	gaa Glu	ggt Gly	att Ile	tat Tyr 95	ttt Phe	288
gca Ala	aaa Lys	aat Asn	ctc Leu 100	acc Thr	cct Pro	gaa Glu	agt Ser	ggt Gly 105	ggt Gly	gcg Ala	att Ile	ggt Gly	tat Tyr 110	gcg Ala	agt Ser	336
ccc Pro	aat Asn	tct Ser 115	cct Pro	acc Thr	gtg Val	gag Glu	att Ile 120	cgt Arg	gat Asp	aca Thr	ata Ile	ggt Gly 125	cct Pro	gta Val	atc Ile	384
ttt Phe	gaa Glu 130	aat Asn	aat Asn	act Thr	tgt Cys	tgc Cys 135	aga Arg	cca Pro	ttt Phe	aca Thr	tcg Ser 140	agt Ser	aat Asn	cct Pro	aat Asn	432
gca Ala 145	gct Ala	gtt Val	aat Asn	aaa Lys	ata Ile 150	aga Arg	gaa Glu	ggc Gly	gga Gly	gcc Ala 155	att Ile	cat His	gct Ala	caa Gln	aat Asn 160	480
ctt Leu	tac Tyr	ata Ile	aat Asn	cat His 165	aat Asn	cat His	gat Asp	gtg Val	gtc Val 170	gga Gly	ttt Phe	atg Met	aag Lys	aac Asn 175	ttt Phe	528
tct Ser	tat Tyr	gtc Val	cga Arg 180	gga Gly	gga Gly	gcc Ala	att Ile	agt Ser 185	acc Thr	gct Ala	aat Asn	acc Thr	ttt Phe 190	gtt Val	gtg Val	576
agc Ser	gag Glu	aat Asn 195	cag Gln	tct Ser	tgt Cys	ttt Phe	ctc Leu 200	ttt Phe	atg Met	gac Asp	aac Asn	atc Ile 205	tgt Cys	att Ile	caa Gln	624
act Thr	aat Asn 210	aca Thr	gca Ala	gga Gly	aaa Lys	ggt Gly 215	ggc Gly	gct Ala	atc Ile	tat Tyr	gct Ala 220	gga Gly	acg Thr	agc Ser	aat Asn	672

FIG. 5A

7969-087 Compared to the ser of the ser of

225	GIU	ser	ASI	230	Cys	Asp	Leu	Pne	235	Asn	Asn	. Ala	240	
		gga Gly							Cys				aat Asn	768
		atc Ile 260												816
		tca Ser												864
		gtt Val												912
		aat Asn												· 960
		ggc Gly												1008
		atc Ile 340												1056
		gct Ala										_		1104
		ggt Gly												1152
		gca Ala									_		_	1200
		tta Leu					Pro							1248
gtc Val		tcc Ser 420												1296
ttt Phe														1344

FIG. 5B

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(SHEET γ OF /4)

						ccc Pro 455										1392
	-	_		_	_	ctt Leu				-						1440
	_	_				aat Asn		_	-	_	_	_				1488
						agc Ser										1536
	_					att Ile	_		_		_					1584
_		_				aat Asn 535		_					-	_		1632
	_					agt Ser										1680
						gaa Glu										1728
	_		_			att Ile			-	_	-		_			1776
	_	_	_	_		tcg Ser				_						1824
						ggc Gly 615										1872
						aca Thr										1920
						tgg Trp										1968
		_	_			ata Ile								_	-	2016

FIG. 5C

ctt Leu	gca Ala	aca Thr 675	gaa Glu	agc Ser	tta Leu	aaa Lys	aat Asn 680	agt Ser	gca Ala	gaa Glu	ctg Leu	aca Thr 685	cct Pro	agt Ser	gat Asp	2064
cat His	cct Pro 690	ttc Phe	tgg Trp	gga Gly	att Ile	aca Thr 695	gga Gly	gga Gly	gga Gly	cta Leu	ggc Gly 700	atg Met	atg Met	gtt Val	tac Tyr	2112
caa Gln 705	gat Asp	cct Pro	cga Arg	gaa Glu	aat Asn 710	cat His	cct Pro	gga Gly	ttc Phe	cat His 715	atg Met	cgc Arg	tct Ser	tcc Ser	gga Gly 720	2160
tac Tyr	tct Ser	gcg Ala	Gly 999	atg Met 725	ata Ile	gca Ala	ggg Gly	cag Gln	aca Thr 730	cac His	acc Thr	ttc Phe	tca Ser	ttg Leu 735	aaa Lys	2208
ttc Phe	agt Ser	cag Gln	acc Thr 740	tac Tyr	acc Thr	aaa Lys	ctc Leu	aat Asn 745	gag Glu	cgt Arg	tac Tyr	gca Ala	aaa Lys 750	aac Asn	aac Asn	2256
gta Val	tct Ser	tct Ser 755	aaa Lys	aat Asn	tac Tyr	tca Ser	tgc Cys 760	caa Gln	gga Gly	gaa Glu	atg Met	ctc Leu 765	ttc Phe	tca Ser	ttg Leu	2304
caa Gln	gaa Glu 770	ggt Gly	ttc Phe	ttg Leu	ctg Leu	act Thr 775	aaa Lys	tta Leu	gtt Val	gly ggg	ctt Leu 780	tac Tyr	agc Ser	tat Tyr	gga Gly	2352
Asp 785	His	Asn	Cys	His	His 790	Phe	Tyr	Thr	caa Gln	Gly .795	Glu	Asn	Leu	Thr	Ser 800	2400
Gln	Gly	Thr	Phe	Arg 805	Ser	Gln	Thr	Met	gga Gly 810	Gly	Ala	Val	Phe	Phe 815	Asp	2448
Leu	Pro	Met	Lys 820	Pro	Phe	Gly	Ser	Thr 825	cat His	Ile	Leu	Thr	Ala 830	Pro	Phe	2496
Leu	Gly	Ala 835	Leu	Gly	Ile	Tyr	Ser 840	Ser	ctg Leu	Ser	His	Phe 845	Thr	Glu	Val	2544
Gly	Ala 850	Tyr	Pro	Arg	Ser	Phe 855	Ser	Thr	aag Lys	Thr	Pro 860	Leu	Ile	Asn	Val	2592
Leu 865	Val	Pro	Ile	Gly	Val 870	Lys	Gly	Ser	ttt Phe	Met 875	Asn	Ala	Thr	Gln	Arg 880	2640
cct Pro	caa Gln	gcc Ala	tgg Trp	act Thr 885	gta Val	gaa Glu	ttg Leu	gca Ala	tac Tyr 890	caa Gln	ccc Pro	gtt Val	ctg Leu	tat Tyr 895	aga Arg	2688

FIG. 5D

gaa Glu						_		_	_	_				2736
ggt Gly	_		_			_	_	_	_					2784
cag Gln 930					_	_					_	tat Tyr		2832
gga Gly												_	-	2880
gct Ala		_		tag										2898

FIG. 5E

					atg Met											48
					ctc Leu											96
					cct Pro											144
tgt Cys	acc Thr 50	atg Met	gag Glu	agc Ser	ctc Leu	ttt Phe 55	cct Pro	gct Ala	ctt Leu	tgt Cys	gct Ala 60	cat His	gca Ala	tca Ser	caa Gln	192
					gta Val 70											240
					gac Asp											288
					aac Asn											336
					cct Pro											384
tcc Ser	ttg Leu 130	cgc Arg	aat Asn	aat Asn	ggt Gly	agc Ser 135	atg Met	agt Ser	ttc Phe	tgt Cys	cga Arg 140	aat Asn	cat His	gct Ala	gaa Glu	432
ggc Gly 145	tct Ser	gga Gly	gga Gly	gcc Ala	atc Ile 150	tct Ser	gcg Ala	gat Asp	gcc Ala	ttt Phe 155	tct Ser	cta Leu	cag Gln	cac His	aac Asn 160	480
tat Tyr	ctt Leu	ttc Phe	aca Thr	gct Ala 165	ttt Phe	gaa Glu	gag Glu	aat Asn	tct Ser 170	tct Ser	aaa Lys	gga Gly	aat Asn	ggc Gly 175	gga Gly	528
gcc Ala	att Ile	cag Gln	gct Ala 180	caa Gln	acc Thr	ttc Phe	tct Ser	tta Leu 185	tct Ser	aga Arg	aat Asn	gtg. Val	tcg Ser 190	cct Pro	att Ile	576
					cgt Arg											624
agt Ser	aat Asn 210	ctt Leu	att Ile	tgt Cys	tca Ser	999 Gly 215	aat Asn	gta Val	aac Asn	cct Pro	ctc Leu 220	ttt Phe	ttc Phe	act Thr	gga Gly	672

FIG. 6A

aac Asn 225	Ser	gcc	acg Thr	aat Asn	gga Gly 230	ggc	gct Ala	att Ile	tgt Cys	tgt Cys 235	atc Ile	agc Ser	gat Asp	cta Leu	aac Asn 240	720
										Cys					cta Leu	768
ttt Phe	gca Ala	agc Ser	aat Asn 260	tct Ser	gct Ala	aaa Lys	gaa Glu	aaa Lys 265	ggc Gly	Gly 399	gct Ala	att Ile	tat Tyr 270	gcc Ala	aag Lys	816
cac His	atg Met	gta Val 275	ttg Leu	cgt Arg	tat Tyr	aac Asn	ggt Gly 280	cct Pro	gtt Val	tcc Ser	ttc Phe	att Ile 285	aac Asn	aac Asn	agc Ser	864
			ggt Gly													912
			ggt Gly													960
			caa Gln													1008
			tct Ser 340													1056
			gta Val								_					1104
			gcc Ala													1152
tta Leu 385	gtt Val	att Ile	cag Gln	Thr	agt Ser 390	gca Ala	aac Asn	cgt Arg	tca Ser	gtg Val 395	att Ile	ttc Phe	tcg Ser	agc Ser	gaa Glu 400	1200
			gaa Glu													1248
cag Gln	cag Gln	Pro	atc Ile 420	gaa Glu	ctg Leu	aaa Lys	Ser	gga Gly 425	cgc Arg	tta Leu	gtt Val	tta Leu	aaa Lys 430	gat Asp	cgc Arg	1296
gct Ala	Val					Ser										1344

FIG. 6B

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	atg Met 450															1392
	acg Thr															1440
	atc Ile															1488
	gga Gly															1536
	aac Asn															1584
cat His	ctt Leu 530	cct Pro	gat Asp	Gly 393	aac Asn	ctc Leu 535	tct Ser	tct Ser	cac His	ttt Phe	gga Gly 540	tat Tyr	caa Gln	gga Gly	gát Asp	1 632
	act Thr															1680
	tgg Trp		Pro													1728
	gtt Val															1776
cag Gln	tcg Ser	atg Met 595	att Ile	aat Asn	aca Thr	ata Ile	gcg Ala 600	cac His	gga Gly	gga Gly	gcc Ala	tat Tyr 605	cta Leu	ttt Phe	gga Gly	1824
acg Thr	tgg Trp 610	gga Gly	tct Ser	gct Ala	gtt Val	tct Ser 615	aat Asn	tta Leu	ttc Phe	tat Tyr	gct Ala 620	cac His	gac Asp	agc Ser	tct Ser	1872
999 Gly 625	aaa Lys	cct Pro	atc Ile	gat Asp	aat Asn 630	tgg Trp	cat His	cat His	aga Arg	agc Ser 635	ctt Leu	ggc Gly	tac Tyr	cta Leu	ttc Phe 640	1920
ggt Gly	atc Ile	agt Ser	act Thr	cac His 645	agt Ser	tta Leu	gat Asp	gac Asp	cat His 650	tct Ser	ttc Phe	tgc Cys	ttg Leu	gct Ala 655	gca Ala	1968
gga Gly	caa Gln	tta Leu	ctc Leu 660	gjå aaa	aaa Lys	tcg Ser	tcc Ser	gat Asp 665	tcc Ser	ttt Phe	att Ile	acg Thr	tct Ser 670	aca Thr	gaa Glu	2016

FIG. 6C

						act Thr											2064
						gca Ala 695											2112
						ttc Phe											2160
						gaa Glu											2208
						agc Ser										-	2256
						gac Asp											2304
						tct Ser 775											2352
						tcc Ser											2400
						gac Asp											2448
						gcc Ala											2496
ttg Leu	gat Asp	tca Ser 835	cga Arg	gcc Ala	tac Tyr	atg Met	ttc Phe 840	agg Arg	ctt Leu	acg Thr	aat Asn	caa Gln 845	aga Arg	gct Ala	cta Leu		2544
cac His	aga Arg 850	ctt Leu	cag Gln	acg Thr	ctg Leu	tta Leu 855	aat Asn	gtg Val	tct Ser	tgt Cys	gtg Val 860	ctg Leu	cgt Arg	gly aaa	caa Gln		2592
agc Ser 865	cat His	agt Ser	tac Tyr	tcc Ser	ctg Leu 870	gat Asp	ctg Leu	gly ggg	acc Thr	act Thr 875	tac Tyr	agg Arg	ttc Phe				2634
tag																	2637

FIG. 6D

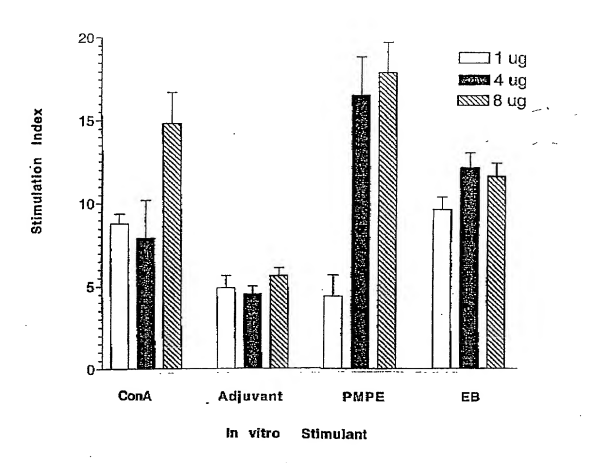


FIG. 7

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DECLARATION AND POWER OF ATTORNEY

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below at 201 et seq. underneath my name.

I believe I am the original, first and sole inventor if only one name is listed at 201 below, or an original, first and joint inventor if plural names are listed at 201 et seq. below, of the subject matter which is claimed and for which a patent is sought on the invention entitled

CHLAMYDIA PROTEIN, GENE SEQUENCE AND USES THEREOF

and for which a patent application:

is attached hereto and includes amendment(s) filed on (grapplicable)

was filed in the United States on as Application No. (for declaration and accompanying application)

with amendment(s) filed on (gapplicable)

O was filed as PCT international Application No. on and was amended under PCT Article 19 on (grapplicable)

I hereby state that I have reviewed and understand the contents of the above identified application, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information known to me to be material to patentability as defined in Title 37, Code of Federal Regulations, §1.56.

I hereby claim foreign priority benefits under Title 35, United States Code, §119(a)-(d) of any foreign application(s) for patent or inventor's certificate listed below and have also identified below any foreign application for patent or inventor's certificate having a filing date before that of the application on which priority is claimed:

EARLIEST FOREIGN APPLIC	'ATION(S), IF ANY, FILED PRIOR '	TO THE FILING DATE C	F THE APPLICATION
APPLICATION NUMBER	COUNTRY	DATE OF FILING (day, month, year)	PRIORITY CLAIMED
			YES □ NO □
			YES 🗆 NO 🗅

I hereby claim the benefit under Title 35, United States Code, §119(e) of any United States provisional application(s) listed below.

APPLICATION NUMBER	FILING DATE

I hereby claim the benefit under Title 35, United States Code, § 120 of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code § 112, I acknowledge the duty to disclose information which is material to patentability as defined in Title 37, Code of Federal Regulations, § 1.56 which became available between the filing date of the prior application and the national or PCT international filing date of this application:

			STATUS	
APPLICATION SERIAL NO.	FILING DATE	PATENTED PATENTED	PENDING	ABANDONED

POWER OF ATTORNEY: As a named inventor, I hereby appoint S. Leslie Misrock (Reg. No. 18872), Berj A. Terzian (Reg. No. 20060), David Weild, III (Reg. No. 21094), Jonathan A. Marshall (Reg. No. 24614), Barry D. Rein (Reg. No. 22411), Stanton T. Lawrence, III (Reg. No. 25736), Charles E. McKenney (Reg. No. 22795), Philip T. Shannon (Reg. No. 24278), Francis E. Morris (Reg. No. 24615), Charles E. Miller (Reg. No. 24576), Gidon D. Stern (Reg. No. 24746), John L. Lauter, Jr. (Reg. No. 27814), Brian M. Foissant (Reg. No. 28462), Brian D. Coggio (Reg. No. 7.624), Rory J. Radding (Reg. No. 28749), Stephen J. Harbulak (Reg. No. 29166), Donald J. Goodell (Reg. No. 19766), James N. Palık (Reg. No. 25510), Thomas E. Friebel (Reg. No. 29258), Laura A. Coruzzi (Reg. No. 30742), Jennier Gordon (Reg. No. 30753), Allan A. Fanucci (Reg. No. 30256), Geraldine F. Baldwin (Reg. No. 31232), Victor N. Balancia (Reg. No. 31231), Samuel B. Abrams (Reg. No. 30605), Steven I. Wallach (Reg. No. 35402), Marcia H. Sundeen (Reg. No. 30893), Paul J. Zegger (Reg. No. 33821), Edmond R. Bannon (Reg. No. 35402), Briuce J. Barker (Reg. No. 33291), Adriane M. Antler (Reg. No. 32605), Thomas G. Rowan (Reg. No. 34419), James G. Markey (Reg. No. 31636), Thomas D. Kohler (Reg. No. 32797), Sont D. Stimpson (Reg. No. 33607), Gary S. Williams (Reg. No. 31066), William S. Galliani (Reg. No. 33885), Ann L. Gisolfi (Reg. No. 31956), Todd A. Wagner (Reg. No. 35299), Scott B. Familant (Reg. No. 35340), Brian D. Siff (Reg. No. 35679), and Alan Tenerbaum (Reg. No. 34939), all of Pennie & Edmonds LLP, whose addresses are 1155 Avenue of the Americas, New York, New York 10036, 1667 K Street N.W., Washington, DC 20006 and 3300 Hallvicw Avenue, Palo Alto, CA 94304, and each of them, my uttorneys, to prosecute this application, and to transact all business in the Petent and Trademark Office connected therewith.

The first fi

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	POST OFFICE ADDRESS	stanet 1687 Armistice Way	cny Marriottsville	STATE OR COUNTRY Maryland	ZIP CODE 21104
	FULL NAME OF INVENTOR	LAST NAME	FIRST NAMB	MIDDLENAME	
0 2	RESIDENCE & CITIZENSHIP	CRY	STATE OR PORBION COUNTRY	COUNTRY OF CITIZENSHIP	
Ī	POST OFFICE ADDRESS	STREET	CHY	STATE OR COMMINY	ZIP CODE
	FULL NAME OF INVENTOR	LAST NAME	FIRET NAME	MIDDLE NAMS	
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	POST OFFICE ADDRESS	STREET	спу	STATE OR COUNTRY	ZIP CODE
	FULL NAME OF INVENTOR	LAST NAME	FIRST NAME	MIDOLENAME	
2 0 4	RESIDENCE & CITIZENSHIP	СПУ	STATE OR FOREIGN COUNTRY	COUNTRY OF CITIZENSHIP	
	POST OFFICE ADDRESS	STREET	CITY	STATICOR COUNTRY	ZIP CODE
	FULL NAME OF INVENTOR	LAST NAME	FIRST NAME	PHITHT LAWR	
2 0 5	RESIDENCE & CITIZENSHIP	CITY	STATE OR FOREIGN COUNTRY	COUNTRY OF CATARASTIP	
	POST OFFICE ADDRESS	Trans	CITY	STATE OR COUNTRY	VIA COLDR
	FULL NAME OF INVENTOR	LASI NAMB	HILET NAME	MIMAN RADDIM	
2 0 6	RESIDENCE &	CITY	STATE OR FOREIGN CCAINTRY	COLINTRY OF CITIZANSHIP	
	POST OFFICE ADDRESS	STRUUT	CITY	STATE OR COUNTRY	ZIP COIDS

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

Section four of this for the Online States Course	SIGNATURE OF INVENTOR 202	SIGNATURE OF INVENTOR 203
SIGNATURE OF INVENTOR 201	and the control of th	
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W. Jarries Jackson		<u> </u>
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SIGNATURE OF INVENTOR 204	SIGNATURE OF ENVENTOR 201	SIGNATURE OF INVENTOR 226
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DATE	P-734.M	

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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Application of: W. James Jackson

Application No.: To be assigned Group Art Unit: To be assigned

Filed: Concurrently herewith Examiner: To be assigned

For: CHLAMYDIA PROTEIN, GENE Attorney Docket No.: 7969-087

SEQUENCE AND USES THEREOF

TRANSMITTAL OF SEQUENCE LISTING UNDER 37 C.F.R. § 1.821

Assistant Commissioner for Patents Washington, D.C. 20231

Sir:

In accordance with 37 C.F.R. § 1.821, Applicant, in connection with the above-identified patent application, submits herewith a Sequence Listing in paper and computer readable form pursuant to 37 C.F.R. §§ 1.821(c) and (e).

I hereby state that the content of the paper and computer readable copies of the Sequence Listing, submitted in accordance with 37 C.F.R. §§ 1.821(c) and (e), respectively, are the same.

Respectfully submitted,

Date October 2, 2000

Geraldine F. Baldwin

(Reg. No.)

PENNIE & EDMONDS LLP 1155 Avenue of the Americas New York, New York 10036-2711

(212) 790-9090

Enclosure

y Nargaret B. Horidalan Reg No. 40, 922

SEQUENCE LISTING

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	< 1	.60>	71													
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cca Pro	gat Asp	cct Pro 35	acg Thr	aaa Lys	gag Glu	tcg Ser	cta Leu 40	tca Ser	aat Asn	aaa Lys	att Ile	agt Ser 45	ttg Leu	aca Thr	gga Gly	144
			aat Asn													192
			cta Leu													240
			cta Leu													288
			ctc Leu 100													336
			cct Pro													384
ttt Phe	gaa Glu 130	aat Asn	aat Asn	act Thr	tgt Cys	tgc Cys 135	aga Arg	cca Pro	ttt Phe	aca Thr	tcg Ser 140	agt Ser	aat Asn	cct Pro	aat Asn	432

	gct Ala															480
	tac Tyr															528
	tat Tyr	_				_		_		_				_	_	576
	gag Glu															624
	aat Asn 210															672
	ttt Phe		_			_	_							_	_	720
	gca Ala															768
_	ggt Gly			_					_	_				_	_	816
	gct Ala															864
_	cta Leu 290	-	_				_						_	_		912
atc Ile 305	aca Thr	aaa Lys	aat Asn	tat Tyr	ggc Gly 310	gga Gly	gct Ala	att Ile	tac Tyr	gct Ala 315	cct Pro	gta Val	gtt Val	acc Thr	cta Leu 320	960
	gat Asp															1008
	ggc															1056
_	cgc Arg		_					-						_		1104

	_							_				_	_	aat Asn	_	1152
		_	_	_				-					_	Gly ggg	_	1200
														gca Ala 415		1248
														gta Val		1296
			_		_			_	_				-	aat Asn		1344
														cta Leu		1392
	_	_		_	_					_				act Thr		1440
	_	-						-	_	_	_	_		aaa Lys 495		1488
	_				_	_		_				_	_	cat His		1536
														cct Pro		1584
														gat Asp		1632
-	_					_	_	_						gat Asp	_	1680
						_			_				_	ctg Leu 575		1728
	_		_						-	_	_		_	cta Leu	_	1776

						tcg Ser										18	824
						ggc Gly 615										18	872
						aca Thr										19	920
aga Arg	acc Thr	tta Leu	tta Leu	ctg Leu 645	act Thr	tgg Trp	ctt Leu	cct Pro	gct Ala 650	ggg Gly	tat Tyr	gtt Val	cct Pro	agc Ser 655	ccg Pro	19	968
						ata Ile										20	016
						aaa Lys										20	064
						aca Thr 695										21	112
						cat His										21	160
Tyr	Ser	Ala	Gly	Met 725	Ile	gca Ala	Gly	Gln	Thr 730	His	Thr	Phe	Ser	Leu 735	Lys	22	802
						aaa Lys										22	256
gta Val	tct Ser	tct Ser 755	aaa Lys	aat Asn	tac Tyr	tca Ser	tgc Cys 760	caa Gln	gga Gly	gaa Glu	atg Met	ctc Leu 765	ttc Phe	tca Ser	ttg Leu	23	304
caa Gln	gaa Glu 770	ggt Gly	ttc Phe	ttg Leu	ctg Leu	act Thr 775	aaa Lys	tta Leu	gtt Val	gly aaa	ctt Leu 780	tac Tyr	agc Ser	tat Tyr	gga Gly	23	352
						ttc Phe										24	100
caa Gln	G1Y 999	acg Thr	ttc Phe	cgt Arg 805	agt Ser	caa Gln	acg Thr	atg Met	gga Gly 810	ggt Gly	gct Ala	gtt Val	ttt Phe	ttt Phe 815	gat Asp	24	48

ctc cct Leu Pro	Met						_							2496
tta ggt Leu Gly	_						_	_						2544
gga gco Gly Ala 850	Tyr													2592
cta gto Leu Val 865				_			_		_		_		-	2640
cct caa Pro Gli	_			_	_	_	_				-			2688
caa gaa Gln Glu	ı Pro					_			_	_	_			2736
ttt ggt Phe Gly	_		_			_	_		_	_				2784
tca cag Ser Gli 930	Gln				_	_						_		2832
cat gga His Gly 945								_					_	2880
att gct Ile Ala	_	_		tag										2898

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<213> Chlamydia sp

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 Gly
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 Ser
 Leu
 Ser
 Gly

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 Arg
 Glu
 Val
 Pro
 Ser
 Arg
 Ile
 Phe
 Leu
 Met
 Pro
 Asn
 Ser
 Val

 Pro
 Asp
 Pro
 Thr
 Lys
 Glu
 Ser
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 Asn
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 Thr
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Ala Ala Thr Phe Ser Leu Ser Asp Val Lys Leu Ser Leu Ile Asp Asp 550 555 Tyr Gly Asn Ser Pro Tyr Glu Ser Thr Asp Leu Thr His Ala Leu Ser 570 Ser Gln Pro Met Leu Ser Ile Ser Glu Ala Ser Asp Asn Gln Leu Arg 585 Ser Asp Asp Met Asp Phe Ser Gly Leu Asn Val Pro His Tyr Gly Trp 600 Gln Gly Leu Trp Thr Trp Gly Trp Ala Lys Thr Gln Asp Pro Glu Pro 620 615 Ala Ser Ser Ala Thr Ile Thr Asp Pro Gln Lys Ala Asn Arg Phe His 630 635 Arg Thr Leu Leu Leu Thr Trp Leu Pro Ala Gly Tyr Val Pro Ser Pro 650 645 Lys His Arg Ser Pro Leu Ile Ala Asn Thr Leu Trp Gly Asn Met Leu 665 Leu Ala Thr Glu Ser Leu Lys Asn Ser Ala Glu Leu Thr Pro Ser Asp 680 His Pro Phe Trp Gly Ile Thr Gly Gly Gly Leu Gly Met Met Val Tyr 695 700 Gln Asp Pro Arg Glu Asn His Pro Gly Phe His Met Arg Ser Ser Gly 710 715 Tyr Ser Ala Gly Met Ile Ala Gly Gln Thr His Thr Phe Ser Leu Lys 725 730 Phe Ser Gln Thr Tyr Thr Lys Leu Asn Glu Arg Tyr Ala Lys Asn Asn 740 745 Val Ser Ser Lys Asn Tyr Ser Cys Gln Gly Glu Met Leu Phe Ser Leu 760 Gln Glu Gly Phe Leu Leu Thr Lys Leu Val Gly Leu Tyr Ser Tyr Gly 775 Asp His Asn Cys His His Phe Tyr Thr Gln Gly Glu Asn Leu Thr Ser 790 795 Gln Gly Thr Phe Arg Ser Gln Thr Met Gly Gly Ala Val Phe Phe Asp 810 Leu Pro Met Lys Pro Phe Gly Ser Thr His Ile Leu Thr Ala Pro Phe 820 825 Leu Gly Ala Leu Gly Ile Tyr Ser Ser Leu Ser His Phe Thr Glu Val 840 Gly Ala Tyr Pro Arg Ser Phe Ser Thr Lys Thr Pro Leu Ile Asn Val 855 860 Leu Val Pro Ile Gly Val Lys Gly Ser Phe Met Asn Ala Thr Gln Arg 870 875 Pro Gln Ala Trp Thr Val Glu Leu Ala Tyr Gln Pro Val Leu Tyr Arg 885 890 Gln Glu Pro Gly Ile Ala Thr Gln Leu Leu Ala Ser Lys Gly Ile Trp 905 900 910 Phe Gly Ser Gly Ser Pro Ser Ser Arg His Ala Met Ser Tyr Lys Ile 920 Ser Gln Gln Thr Gln Pro Leu Ser Trp Leu Thr Leu His Phe Gln Tyr 935 940 His Gly Phe Tyr Ser Ser Ser Thr Phe Cys Asn Tyr Leu Asn Gly Glu 950 955 Ile Ala Leu Arg Phe

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tca Ser	tco Ser	aca Thr	gcg Ala 20	Val	ctc Leu	ttt Phe	ggc	cag Gln 25	Asp	ccc Pro	tta Leu	ı ggt ı Gly	gaa Glu 30	Thr	gcc Ala	96
ctc Leu	cto Leu	act Thr	. Ta	aat Asn	cct Pro	aat Asn	cat His 40	Val	gto Val	tgt Cys	aca Thr	ttt Phe 45	Phe	gag Glu	gac Asp	144
tgt Cys	acc Thr 50	Met	gag Glu	agc Ser	ctc Leu	ttt Phe 55	cct Pro	gct Ala	ctt Leu	tgt Cys	gct Ala 60	His	gca Ala	tca Ser	caa Gln	192
gac Asp 65	gat Asp	cct Pro	ttg Leu	tat Tyr	gta Val 70	ctt Leu	gga Gly	aat Asn	tcc Ser	tac Tyr 75	tgt Cys	tgg Trp	ttc Phe	gta Val	tct Ser 80	240
aaa Lys	ctc Leu	cat His	atc Ile	acg Thr 85	gac Asp	ccc Pro	aaa Lys	gag Glu	gct Ala 90	ctt Leu	ttt Phe	aaa Lys	gaa Glu	aaa Lys 95	gga Gly	288
gat Asp	ctt Leu	tcc Ser	att Ile 100	caa Gln	aac Asn	ttt Phe	cgc Arg	ttc Phe 105	ctt Leu	tcc Ser	ttc Phe	aca Thr	gat Asp 110	tgc Cys	tct Ser	336
tcc Ser	aag Lys	gaa Glu 115	agc Ser	tct Ser	cct Pro	tct Ser	att Ile 120	att Ile	cat His	caa Gln	aag Lys	aat Asn 125	ggt Gly	cag Gln	tta Leu	384
tcc Ser	ttg Leu 130	cgc Arg	aat Asn	aat Asn	ggt Gly	agc Ser 135	atg Met	agt Ser	ttc Phe	tgt Cys	cga Arg 140	aat Asn	cat His	gct Ala	gaa Glu	432
ggc Gly 145	tct Ser	gga Gly	gga Gly	gcc Ala	atc Ile 150	tct Ser	gcg Ala	gat Asp	gcc Ala	ttt Phe 155	tct Ser	cta Leu	cag Gln	cac His	aac Asn 160	480
tat Tyr	ctt Leu	ttc Phe	aca Thr	gct Ala 165	ttt Phe	gaa Glu	gag Glu	aat Asn	tct Ser 170	tct Ser	aaa Lys	gga Gly	aat Asn	ggc Gly 175	gga Gly	528
gcc Ala	att Ile	cag Gln	gct Ala 180	caa Gln	acc Thr	ttc Phe	tct Ser	tta Leu 185	tct Ser	aga Arg	aat Asn	gtg Val	tcg Ser 190	cct Pro	att Ile	576
tct Ser	ttc Phe	gcc Ala 195	cgt Arg	aat Asn	cgt Arg	Ala	gat Asp 200	tta Leu	aat Asn	ggc Gly	ggc	gct Ala 205	att Ile	tgc Cys	tgt Cys	624

agt Sei	aat Asr 210	ı Leı	att Ile	tgt Cys	tca Ser	999 Gly 215	aat Asr	gta Val	aac Asn	cct Pro	cto Leu 220	Phe	ttc Phe	act Thr	gga Gly	672
aac Asr 225	ı Ser	gcc Ala	acg Thr	aat Asn	gga Gly 230	Gly	gct Ala	att Ile	tgt Cys	tgt Cys 235	Ile	ago Ser	gat Asp	cta Leu	aac Asn 240	720
acc Thr	tca Ser	gaa Glu	aaa Lys	ggc Gly 245	Ser	ctc Leu	tct Ser	ctt Leu	gct Ala 250	tgt Cys	aac Asn	caa Gln	gaa Glu	acg Thr 255	cta Leu	768
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cac His	atg Met	gta Val 275	ttg Leu	cgt Arg	tat Tyr	aac Asn	ggt Gly 280	cct Pro	gtt Val	tcc Ser	ttc Phe	att Ile 285	aac Asn	aac Asn	agc Ser	864
gct Ala	aaa Lys 290	ata Ile	ggt Gly	gga Gly	gct Ala	atc Ile 295	gcc Ala	atc Ile	cag Gln	tcc Ser	gga Gly 300	gly ggg	agt Ser	ctc Leu	tct Ser	912
atc Ile 305	ctt Leu	gca Ala	ggt Gly	gaa Glu	gga Gly 310	tct Ser	gtt Val	ctg Leu	ttc Phe	cag Gln 315	aat Asn	aac Asn	tcc Ser	caa Gln	cgc Arg 320	960
acc Thr	tcc Ser	gac Asp	caa Gln	ggt Gly 325	cta Leu	gta Val	aga Arg	aac Asn	gcc Ala 330	atc Ile	tac Tyr	tta Leu	gag Glu	aaa Lys 335	gat Asp	1008
gcg Ala	att Ile	ctt Leu	tct Ser 340	tcc Ser	tta Leu	gaa Glu	gct Ala	cgc Arg 345	aac Asn	gga Gly	gat Asp	att Ile	ctt Leu 350	ttc Phe	ttt Phe	1056
gat Asp	cct Pro	att Ile 355	gta Val	caa Gln	gaa Glu	agt Ser	agc Ser 360	agc Ser	aaa Lys	gaa Glu	tcg Ser	cct Pro 365	ctt Leu	ccc Pro	tcc Ser	1104
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tta Leu 385	gtt Val	att Ile	cag Gln	aca Thr	agt Ser 390	gca Ala	aac Asn	cgt Arg	tca Ser	gtg Val 395	att Ile	ttc Phe	tcg Ser	agc Ser	gaa Glu 400	1200
cgt Arg	ctt Leu	tct Ser	gaa Glu	gaa Glu 405	gaa Glu	aaa Lys	act Thr	Pro	gat Asp 410	aac Asn	ctc Leu	act Thr	tcc Ser	caa Gln 415	cta Leu	1248
cag Gln	cag Gln	cct Pro	atc Ile 420	gaa Glu	ctg Leu	aaa Lys	tcc Ser	gga Gly 425	cgc Arg	tta Leu	gtt Val	tta Leu	aaa Lys 430	gat Asp	cgc Arg	1296

gct Ala	gtc Val	ctt Leu 435	tcc Ser	gcg Ala	cct Pro	tct Ser	ctc Leu 440	tct Ser	cag Gln	gat Asp	cct Pro	caa Gln 445	gct Ala	ctc Leu	ctc Leu	1344
					act Thr											1392
					ccc Pro 470											1440
					aac Asn											1488
					ttt Phe											1536
					ctc Leu											1584
cat His	ctt Leu 530	cct Pro	gat Asp	Gly 999	aac Asn	ctc Leu 535	tct Ser	tct Ser	cac His	ttt Phe	gga Gly 540	tat Tyr	caa Gln	gga Gly	gat Asp	1632
					aaa Lys 550											1680
Asn	Trp	Thr	Pro	Lys 565	aac Asn	Tyr	Val	Pro	His 570	Pro	Glu	Arg	Gln	Ser 575	Thr	1728
Leu	Val	Ala	Asn 580	Thr	ctt Leu	Trp	Asn	Thr 585	Tyr	Ser	Asp	Met	Gln 590	Āla	Val	1776
Gln	Ser	Met 595	Ile	Asn	aca Thr	Ile	Ala 600	His	Gly	Gly	Ala	Tyr 605	Leu	Phe	Gly	1824
acg Thr	tgg Trp 610	gga Gly	tct Ser	gct Ala	gtt Val	tct Ser 615	aat Asn	tta Leu	ttc Phe	tat Tyr	gct Ala 620	cac His	gac Asp	agc Ser	tct Ser	1872
Gly 625	Lys	Pro	Ile	Asp	aat Asn 630	Trp	His	His	Arg	Ser 635	Leu	Gly	Tyr	Leu	Phe 640	1920
ggt Gly	atc Ile	agt Ser	act Thr	cac His 645	agt Ser	tta Leu	gat Asp	gac Asp	cat His 650	tct Ser	ttc Phe	tgc Cys	ttg Leu	gct Ala 655	gca Ala	1968

gga Gly	caa Gln	tta Leu	ctc Leu 660	Gly	aaa Lys	tcg Ser	tcc Ser	gat Asp 665	Ser	ttt Phe	att Ile	acg Thr	tct Ser 670	Thr	gaa Glu	2016
acg Thr	acc Thr	tcc Ser 675	tat Tyr	ata Ile	gct Ala	act Thr	gta Val 680	caa Gln	gcg Ala	caa Gln	ctc Leu	gct Ala 685	acc Thr	tct Ser	cta Leu	2064
atg Met	aaa Lys 690	atc Ile	tct Ser	gca Ala	cag Gln	gca Ala 695	tgc Cys	tac Tyr	aat Asn	gaa Glu	agt Ser 700	Ile	cat His	gag Glu	cta Leu	2112
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aat Asn	ggt Gly	tcc Ser	gga Gly 740	ctg Leu	ttc Phe	agc Ser	tcc Ser	ttc Phe 745	tct Ser	att Ile	ttc Phe	tct Ser	aaa Lys 750	ctg Leu	caa Gln	2256
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cgg Arg	tcc Ser 770	ttt Phe	tct Ser	gcc Ala	agc Ser	tct Ser 775	ttc Phe	aga Arg	aat Asn	att Ile	tca Ser 780	ctt Leu	cct Pro	ata Ile	gga Gly	2352
ata Ile 785	aca Thr	ttt Phe	gaa Glu	aaa Lys	aaa Lys 790	tcc Ser	caa Gln	aaa Lys	aca Thr	cga Arg 795	acc Thr	tac Tyr	tat Tyr	tac Tyr	ttt Phe 800	2400
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cac His	aga Arg 850	ctt Leu	cag Gln	acg Thr	ctg Leu	tta Leu 855	aat Asn	gtg Val	tct Ser	tgt Cys	gtg Val 860	ctg Leu	cgt Arg	gly aaa	caa Gln	2592
agc Ser 865	cat His	agt Ser	tac Tyr	tcc Ser	ctg Leu 870	gat Asp	ctg Leu	gl ^à aaa	acc Thr	act Thr 875	tac Tyr	agg Arg	ttc Phe			2634
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Gln Gln Pro Ile Glu Leu Lys Ser Gly Arg Leu Val Leu Lys Asp Arg Ala Val Leu Ser Ala Pro Ser Leu Ser Gln Asp Pro Gln Ala Leu Leu Ile Met Glu Ala Gly Thr Ser Leu Lys Thr Ser Ser Asp Leu Lys Leu Ala Thr Leu Ser Ile Pro Leu His Ser Leu Asp Thr Glu Lys Ser Val Thr Ile His Ala Pro Asn Leu Ser Ile Gln Lys Ile Phe Leu Ser Asn Ser Gly Asp Glu Asn Phe Tyr Glu Asn Val Glu Leu Leu Ser Lys Glu Gln Asn Asn Ile Pro Leu Leu Thr Leu Ser Lys Glu Gln Ser His Leu His Leu Pro Asp Gly Asn Leu Ser Ser His Phe Gly Tyr Gln Gly Asp Trp Thr Phe Ser Trp Lys Asp Ser Asp Glu Gly His Ser Leu Ile Ala Asn Trp Thr Pro Lys Asn Tyr Val Pro His Pro Glu Arg Gln Ser Thr Leu Val Ala Asn Thr Leu Trp Asn Thr Tyr Ser Asp Met Gln Ala Val Gln Ser Met Ile Asn Thr Ile Ala His Gly Gly Ala Tyr Leu Phe Gly Thr Trp Gly Ser Ala Val Ser Asn Leu Phe Tyr Ala His Asp Ser Ser Gly Lys Pro Ile Asp Asn Trp His His Arg Ser Leu Gly Tyr Leu Phe Gly Ile Ser Thr His Ser Leu Asp Asp His Ser Phe Cys Leu Ala Ala Gly Gln Leu Leu Gly Lys Ser Ser Asp Ser Phe Ile Thr Ser Thr Glu Thr Thr Ser Tyr Ile Ala Thr Val Gln Ala Gln Leu Ala Thr Ser Leu Met Lys Ile Ser Ala Gln Ala Cys Tyr Asn Glu Ser Ile His Glu Leu Lys Thr Lys Tyr Arg Ser Phe Ser Lys Glu Gly Phe Gly Ser Trp His Ser Val Ala Val Ser Gly Glu Val Cys Ala Ser Ile Pro Ile Val Ser Asn Gly Ser Gly Leu Phe Ser Ser Phe Ser Ile Phe Ser Lys Leu Gln Gly Phe Ser Gly Thr Gln Asp Gly Phe Glu Glu Ser Ser Gly Glu Ile Arg Ser Phe Ser Ala Ser Ser Phe Arg Asn Ile Ser Leu Pro Ile Gly Ile Thr Phe Glu Lys Lys Ser Gln Lys Thr Arg Thr Tyr Tyr Phe Leu Gly Ala Tyr Ile Gln Asp Leu Lys Arg Asp Val Glu Ser Gly Pro Val Val Leu Leu Lys Asn Ala Val Ser Trp Asp Ala Pro Met Ala Asn Leu Asp Ser Arg Ala Tyr Met Phe Arg Leu Thr Asn Gln Arg Ala Leu His Arg Leu Gln Thr Leu Leu Asn Val Ser Cys Val Leu Arg Gly Gln Ser His Ser Tyr Ser Leu Asp Leu Gly Thr Thr Tyr Arg Phe

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 Thr Gly Asp Thr His Asn Leu Thr Asn Cys
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Ser Val Pro Asp Pro Thr Lys Glu Ser Leu Ser Asn Lys Ile Ser Leu
                                25
Thr Gly Asp Thr His Asn Leu Thr Asn Cys Tyr Leu Asp Asn Leu Arg
                            40
Tyr Ile Leu Ala Ile Leu Gln Lys Thr Pro Asn Glu Gly Ala Ala Val
                        55
Thr Ile Thr Asp Tyr Leu Ser Phe Phe Asp Thr Gln Lys Glu Gly Ile
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Tyr Phe Ala Lys Asn Leu Thr Pro Glu Ser Gly Gly Ala Ile Gly Tyr
Ala Ser Pro Asn Ser Pro Thr Val Glu Ile Arg
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Leu Arg Tyr Ile Leu Ala Ile Leu Gln Lys Thr Pro Asn Glu Gly Ala
           20
                               25
Ala Val Thr Ile Thr Asp Tyr Leu Ser Phe Phe Asp Thr Gln Lys Glu
                           40
Gly Ile Tyr Phe Ala Lys Asn Leu Thr Pro Glu Ser Gly Gly Ala Ile
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Gly Tyr Ala Ser Pro Asn Ser Pro Thr Val Glu Ile Arg Asp Thr Ile
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Gly
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Met Lys Asn Phe Ser Tyr Val Arg Gly Gly Ala Ile Ser Thr Ala Asn
 Thr Phe
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                                25
Glu Ser Asn Asn Cys Asp Leu Phe Phe Ile Asn Asn Ala Cys Cys Ala
                            40
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Asn Ile Val
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Asp Val Thr Gly Asn Arg Gly Arg Ile Phe Phe Ser Asp Asn Ile Thr
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Lys Asn Tyr Gly Gly Ala Ile Tyr Ala Pro Val Val Thr Leu Val Asp
                            40
Asn Gly Pro Thr Tyr Phe Ile Asn Asn Ile Ala Asn Asn Lys Gly Gly
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                                            60
Ala Ile Tyr Ile Asp Gly Thr Ser Asn Ser Lys Ile Ser Ala Asp Arg
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His Ala Ile Ile Phe Asn Glu Asn Ile Val Thr Asn
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Asn Ser
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Leu Thr Leu Ser Asn Gly Phe Leu Cys Ile Glu Asp His Ala Gln Leu
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Asn Tyr Ser Cys Gln Gly Glu Met Leu Phe Ser Leu Gln Glu Gly Phe
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Asp Thr His Asn Leu Thr Asn Cys Tyr Leu Asp Asn Leu Arg Tyr Ile
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Ala Lys Asn Leu Thr Pro Glu Ser Gly Gly Ala Ile Gly Tyr Ala Ser
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Thr Phe Ser Leu Ser Arg Asn Val Ser Pro Ile Ser Phe Ala Arg Asn
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Leu Ser Glu Glu Glu Lys Thr Pro Asp Asn Leu Thr Ser Gln Leu Gln
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Cys Thr Met Glu Ser Leu Phe Pro Ala Leu Cys Ala His Ala Ser Gln
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Asp Asp Pro Leu Tyr Val Leu Gly Asn Ser Tyr Cys Trp Phe Val Ser
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Lys Leu His Ile Thr Asp Pro Lys Glu Ala Leu Phe Lys Glu Lys Gly
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Asp Leu Ser Ile Gln Asn Phe Arg Phe Leu Ser Phe Thr Asp Cys Ser
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105

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Ala Ile Gln Ala Gln Thr Phe Ser Leu Ser Arg Asn Val Ser Pro Ile
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Ala Lys Ile Gly Gly Ala Ile Ala Ile Gln Ser Gly Gly Ser Leu Ser
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